

JULY 1973

American Heart Journal

International Editorial Board

J. A. Abildskov *Salt Lake City*
 P. P. Ahlquist *Augusta*
 A. Ben-Chimol *Phoenix*
 S. Gilbert Blount Jr. *Denver*
 Yves Bouvrain *Paris*
 Daniel A. Brody *Memphis*
 Agustín Castellanos Jr. *Miami*
 Muir Clapper *Detroit*
 J. M. Criley *Torrance Calif*
 James A. Cronvich *New Orleans*
 Anthony N. Damato *Staten Island N Y*
 Arthur C. DeGraff *New York*
 Lewis Dexter *Boston*
 Leonard S. Dreifus *Philadelphia Pa*
 Charles Dubost *Paris*
 Jesse E. Edwards *St Paul*
 Harvey Feigenbaum *Indianapolis*
 Alvan R. Feinstein *New Haven Conn*
 M. Irénée Ferter *New York*
 Charles Fuch *Indianapolis*
 Julian Frieden *New Rochelle N Y*
 Peter C. Gases *Charleston*
 A. Sidney Harris *New Orleans*
 George M. Hass *Chicago*
 Frantisek Herles *Prague Czechoslovakia*
 Paul Rugenoltz *Rotterdam*
 Mohamed Ibrahim *Cairo Egypt*
 Harold A. Kahn *Bethesda*
 P. I. Korner *Sydney*
 Richard Langendorf *Chicago*
 Maurice Lev *Chicago*
 Robert L. Levy *New York*
 Howard P. Lewis *Portland Ore*
 R. J. Linden *Leeds*
 F. Loogen *Dusseldorf Germany*
 Daniel S. Lukas *New York*
 Pavel E. Lukomsky *Moscow*
 Dwight C. McGoon *Rochester Minn*
 Felipe Mendoza *Mexico D F Mex*
 Gordon K. Moe *Utica N Y*
 Clifford V. Nelson *Portland Me*
 Edward S. Orgain *Durham*
 Victor Parsonnet *Newark*
 Joseph H. Perloff *Philadelphia*
 Alfred Pick *Chicago*
 Simon Rodbard *Duarte*
 Ralph C. Scott *Cincinnati*
 Ewald E. Selkurt *Indianapolis*
 Arthur Selzer *San Francisco*
 J. P. Shillingford *London*
 Ernst Simonson *Minneapolis*
 John R. Smith *St Louis*
 Louis A. Soloff *Philadelphia*
 Madison S. Spaeh *Durham*
 David H. Spodick *Boston*
 Tadasu Takatsu *Osaka*
 Jack L. Titus *Rochester Minn*
 William H. Weidman *Rochester Minn*
 Myron W. Wheat Jr. *Louisville*
 Henry *Cleveland*

GEORGE E BURCH *Editor*

HARRY L COLCOLOUGH

JOHN H PHILLIPS *Assistant Editors*

THE C V MOSBY COMPANY *Publisher*

St Louis Mo 63141 USA

An international
 publication
 for the study of
 the circulation

Contents on pp 3 5 and 7

AHJOA2 86 (1) 1 148 (1973)

American Heart Journal

An international publication for the study of the circulation

Editor

GEORGE E BURCH M D

Assistant editors

HARRY I COLCOLOUGH M D

THOMAS D GILES M D

1430 Tulane Avenue

New Orleans Louisiana 70112

Publisher

THE C V MOSBY COMPANY

11830 Westline Industrial Drive

St Louis Missouri 63141

Editorial communications

Original communications Manuscripts for publication letters and all other communications relating to the editorial management of the Journal should be sent to the Editor Dr George E Burch 1430 Tulane Avenue New Orleans Louisiana 70112. Articles are accepted for publication with the understanding that they are contributed solely to the American Heart Journal.

Neither the editor nor the publisher accepts responsibility for the views and statements of authors whose manuscripts are published as original communications.

Manuscripts Manuscripts must be type written (on one side of the paper only) with liberal margins and completely double spaced except for mathematical material which must be triple spaced. When symbols cannot be inserted by the typewriter they must be clearly identifiable and carefully aligned. A list of References must appear at the end of the article following style used in the Cumulated Index Medicus and giving authors names and initials title and the name volume number page number and year of publication of the journal in that order. Illustrations accompanying manuscripts must be numbered provided with suitable legends and marked lightly on the back with the author's name. Submission of two copies of a manuscript and its illustrations will expedite editorial handling and speed publication.

Illustrations A reasonable number of halftone illustrations will be reproduced free of cost to the author but special arrangements

must be made with the editor for color plates elaborate tables or extra illustrations. To insure clear reproduction all copy for zinc cuts including pen drawings and charts must be prepared with India ink and a black ribbon must be used for type written material. Only good photographic prints and original drawings should be supplied for halftone work.

Exchanges Contributions letters exchanges reprints and all other communications relating to the Journal should be sent to Dr George E Burch 1430 Tulane Avenue New Orleans Louisiana 70112. Writers on subjects which are related in any way to cardiovascular disease are requested to place this address on their permanent reprint mailing lists.

Reprints Reprints of articles must be ordered directly through the publishers The C V Mosby Co 11830 Westline Industrial Drive St Louis Mo 63141 U S A who will quote prices upon publication of the article. Individual reprints of an article must be obtained through the author.

Review of books Publishers and authors are informed that the space of the Journal is so fully occupied by matter pertaining to the branches to which it is devoted that only works treating of these subjects can be noticed. Books and monographs on the anatomy physiology pharmacology therapeutics and pathology of the heart blood vessels and circulation will be reviewed when space is available. Send books to the Editor Dr George E Burch 1430 Tulane Avenue New Orleans Louisiana 70112.

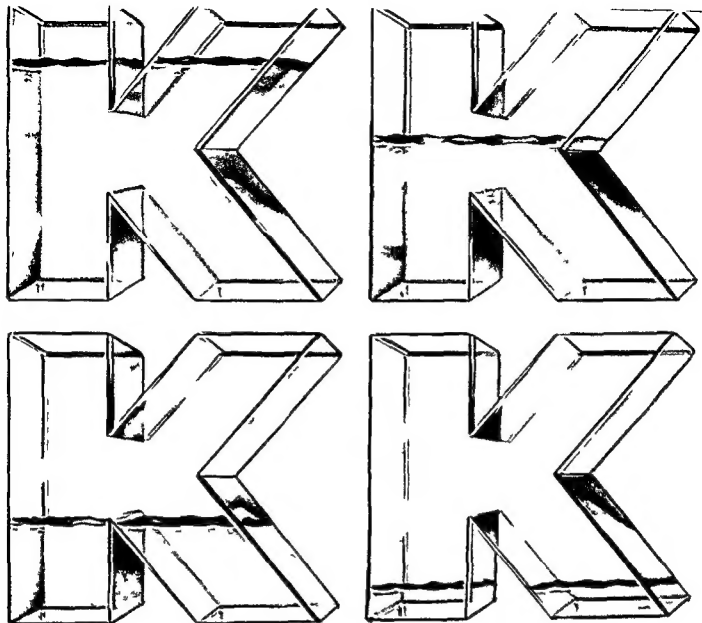
Now Available

A series of brochures from
Cooper Laboratories dealing with
Sudden Coronary Death
To reserve your series, contact your
Cooper representative or write to
Cooper Colloquium,
Cooper Laboratories, Fairfield Road,
Wayne, New Jersey 07470

In the treatment of atrial fibrillation...

it's one thing to restore the rhythm





'Dyrenium' Conserves Potassium add it to the thiazides or furosemide

In the treatment of edema*

Dyrenium® conserves the potassium most other diuretics waste and augments the action of other diuretics

Before prescribing, see complete prescribing information in SK&F literature or PDR

Indications: Edema associated with congestive heart failure, cirrhosis and nephrotic syndrome, steroid-induced edema, idiopathic edema, edema due to secondary hyperaldosteronism and edema resistant to other diuretic therapy

Contraindications: Severe or progressive kidney disease or dysfunction (possible exception: nephrosis); Severe hepatic disease; Pre-existing elevated serum potassium; Hypersensitivity to the drug; Continued use in developing hyperkalemia; Do not give potassium supplements either by drug or by diet

Warnings: Observe regularly for possible blood dyscrasias, liver damage or other

idiosyncratic reactions. Blood dyscrasias have been reported. Check BUN and serum potassium periodically, especially in the elderly, diabetics and those with suspected or confirmed renal insufficiency. Use in pregnancy only when essential to patient welfare.

Dyrenium (triamterene, SK&F) and spironolactone are not usually used concurrently if they are, however, frequent serum potassium determinations are required.

Precautions: If hyperkalemia develops, withdraw the drug. The following may also occur: electrolyte imbalance, low salt syndrome (with low salt intake), reversible mild nitrogen retention, decreasing alkali reserve with possible metabolic acidosis. Do periodic hematologic studies in cirrhotics with splenomegaly. Concomitant use with anti-hypertensive drugs may result in an additive hypotensive effect. When Dyrenium is to be discontinued after intensive or prolonged therapy, withdraw gradually because of possible rebound kaliuresis.

Adverse Reactions: Diarrhea, nausea and vomiting (may indicate electrolyte imbalance); other gastrointestinal disturbances; weakness, headache, dry mouth, anaphylaxis; photosensitivity; elevated uric acid; rash.

Note: When combined with another diuretic, the initial dosage of each agent should be lower than recommended. Supplied 100 mg capsules in bottles of 100.

SK&F Co., Carolina, PR 00630
A subsidiary of
Smith Kline & French Laboratories

100 mg capsules

Dyrenium®
brand of triamterene

Contents

Editorial

Isoproterenol in cardiology 149

R P Ahlquist PhD FCP Augusta Ga

Clinical communications

Changes in the coronary vasculature in endomyocardial fibrosis and their possible significance 152

*Zilton A Andrade MD and Antonio R L Teixeira MD
Bahia Brazil*

Comparison of pulmonary wedge and left atrial pressure in man 159

*Abe Walston II MD and M Eugene Kendall MD
Durham N C*

The progression of coronary atherosclerotic disease as assessed by cine-coronary arteriography 163

*Robert R Henderson MD and George G Rowe MD
Madison Wis*

Bifascicular block. A clinical and electrophysiologic study 173

*Dorothy Kuntzsch MD Manohar Punja MD Norman Cagen MD
Pa Fernando MD Barrie Levitt MD and
Yusuf Z Yucoglu MD FACC New York N Y*

Systolic time intervals in pregnancy and the postpartum period 182

*Shirley Rubler MD FACC Ralph Schneebaum MD and
Nina Hammer BA New York N Y*

Experimental and laboratory reports

Mathematical relationship between automaticity of the sinus node and the AV junction 189

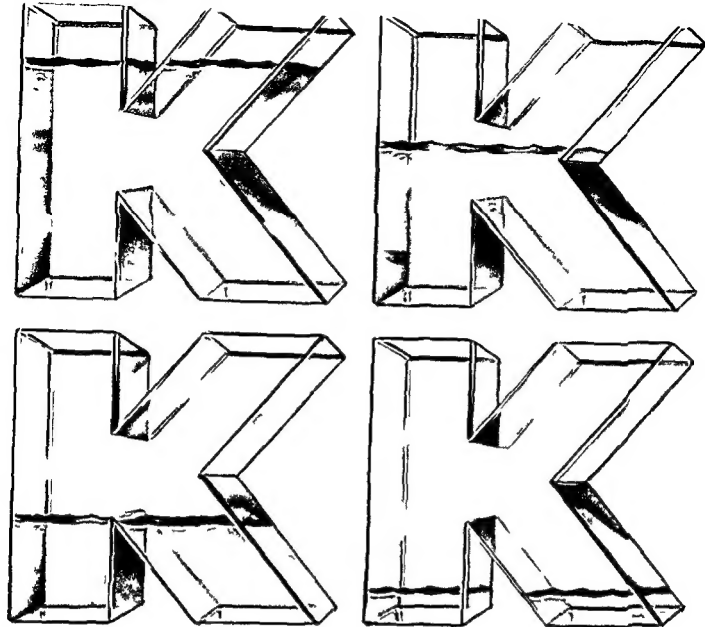
*Ferdinand Lrihaler MD Charles R Katchels PhD Josiah Macy Jr PhD
and Thomas W James MD Birmingham Ala*

Comparative surface potential patterns in obstructive and nonobstructive cardiomyopathy 196

Nancy C Flowers MD and Leo G Horan MD Augusta Ga

The effect of heart rate acetylcholine and vagal stimulation on antegrade and retrograde His Purkinje conduction in the intact heart 203

*P Jacob LaResse MD Anthony N Damato MD Sun H Lau MD
Masood Akhtar MD and Gustavus A Bobb Staten Island N Y*



'Dyrenium' Conserves Potassium add it to the thiazides or furosemide

In the treatment of edema*

Dyrenium conserves the potassium most other diuretics waste and augments the action of other diuretics

Before prescribing see complete prescribing information in SK&F literature or PDR

Indications Edema associated with congestive heart failure, cirrhosis and nephrotic syndrome, steroid-induced edema, idiopathic edema, edema due to secondary hyperaldosteronism and edema resistant to other diuretic therapy

Contraindications Severe or progressive kidney disease or dysfunction (possible exception: nephrosis) Severe hepatic disease Pre-existing elevated serum potassium Hypersensitivity to the drug Continued use in developing hypokalemia Do not give potassium supplements either by drug or by diet

Warnings Observe regularly for possible blood dyscrasias, liver damage or other

idiosyncratic reactions. Blood dyscrasias have been reported. Check BUN and serum potassium periodically especially in the elderly, diabetics and those with suspected or confirmed renal insufficiency. Use in pregnancy only when essential to patient welfare.

Dyrenium (triamterene, SK&F) and spironolactone are not usually used concurrently. If they are, however, frequent serum potassium determinations are required.

Precautions If hypokalemia develops, withdraw the drug. The following may also occur: electrolyte imbalance, low salt syndrome (with low salt intake), reversible mild nitrogen retention, decreasing alkali reserve with possible metabolic acidosis. Do periodic hematologic studies in cirrhotics with splenomegaly. Concomitant use with anti-hypertensive drugs may result in an additive hypotensive effect. When Dyrenium is to be discontinued after intensive or prolonged therapy, withdraw gradually because of possible rebound kaliuresis.

Adverse Reactions Diarrhea, nausea and vomiting (may indicate electrolyte imbalance), other gastrointestinal disturbances, weakness, headache, dry mouth, anaphylaxis, photosensitivity, elevated uric acid, rash.

Note When combined with another diuretic, the initial dosage of each agent should be lower than recommended.

Supplied 100 mg capsules in bottles of 100

SK&F Co., Carolina, P.R. 00630
A subsidiary of
Smith, Kline & French Laboratories

100 mg capsules

Dyrenium®
brand of triamterene

Contents *continued*

Drug failure in reducing pressor effect of isometric handgrip stress test in hypertension 211

Sejjan Lamić MD and Frederick W Wolff MD Washington D C

The role of beta adrenoceptors in the coronary and systemic hemodynamic responses to emotional stress in conscious dogs 216

1/ Bergamaschi Ph D A M Carozzini DVM V Mandelli D Sc
and R G Shanks MD Milan Italy

Case reports

Cardiac lymphangioma and lipoma

Report of a case of simultaneous occurrence in association with lipomatous infiltration of the myocardium and cardiac arrhythmia 227

Daniel T. Anbe M D and Gerald Fine M D Detroit Mich

Propranolol induced alopecia 236

Carroll M Martin MD Major MC USA Edward G Sonnenfeld MD
Major MC USA and Howard I Maibach MD
San Francisco Calif

Clinical pathologic conference

Clinical pathologic conference 238

Yake A Kara horlu M D Relf M Gunnar M D and
Cecil A Krakower M D Chicago Ill

Fundamentals of clinical cardiology

Mechanisms of cardiac arrhythmias From hypothesis to physiologic fact 249

Alfred Pick M D Chicago Ill

Appraisal and reappraisal of cardiac therapy

Drug therapy of heart disease in pediatric patients II

The treatment of congestive heart failure in infants and children with digitalis preparations 270

Wonska M Ruthowski MD Sanford & Cohen MD and
Eugenia F Doyle MD New York N Y

continued on page 7

U.S. GOVERNMENT PRINTING OFFICE: 1973

| | L. 5 | Canada \$100 | Other 0.00 |
|------------|---------|-----------------|---------------|
| Individual | \$26.0 | \$29.50 | \$30.25 |
| Person 1) | \$19.50 | \$22.50 | \$23.25 |
| Individual | \$13.6 | \$16.65 | \$17.40 |

pay ble to this Bureau.

1. All titles and materials are available to public and private libraries, schools, hospitals, and private citizens and organizations for use and deposit in their collections and for personal subscription and all subscription must be in the name of and billed to individuals.

Subscriptions may be at a y tim

Second class post paid at St. Louis Mo.
Printed in the U. S. A.

Printed in the U. S. A. Copyright © 1975 by The C. V. Mosby Company

**“The drug of choice
for oral replacement
of potassium is
potassium chloride
solution.”**

AMA Drug Evaluations 1971 First Edition
Chicago: American Medical Association p. 121

**Kay Ciel® Elixir is
potassium chloride...
tastes good too!**

COMPOSITION: Each 15 cc. (one table spoonful) contains potassium chloride 15 Gm. supplying 20 mEq. of elemental potassium in a cherry flavored palatable base alcohol 4. Contains no sugar.

INDICATIONS: Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy; digitalis intoxication; low dietary intake of potassium or as a result of excessive vomiting and diarrhea.

CONTRAINDICATIONS: Impaired renal function; untreated Addison's Disease; dehydration; heat cramps; and hyperkalemia.

PRECAUTIONS: Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient, since the amount of deficiency and corresponding daily dose is

often not known. Excessive or even therapeutic dosages may result in potassium intoxication. Patients should be frequently checked and periodic ECG and/or plasma potassium levels made. High plasma concentrations of potassium ion may cause cardiac depression, arrhythmias or arrest. Use with caution in patients with cardiac disease. In hypokalemic states attention should be directed toward the correction of the frequently associated hypochloremic alkalosis.

SIDE EFFECTS: Vomiting, nausea, abdominal discomfort and diarrhea may occur. Symptoms and signs of potassium intoxication include listlessness, mental confusion, paresthesia of the extremities, weakness of the legs, flaccid paralysis, fall in blood pressure, cardiac arrhythmias and heart block. When hyperkalemia

exists, it should be promptly treated with the discontinuance of potassium administration or other steps to lower serum levels. If indicated, since sudden shift in plasma levels may induce potentially dangerous cardiac arrhythmias.

DOSEAGE AND ADMINISTRATION: Adults: one tablespoonful (15 cc.) diluted in one glass of water, twice daily after the morning and evening meal. Larger doses may be indicated according to the individual patient's requirements but should be administered under close supervision due to the possibility of potassium intoxication. Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal irritation.
HOW SUPPLIED: One pint and one gallon bottles.

Cooper

Cooper Laboratories Inc. Wayne, N.J. 07470/St. Therese, P.Q. Canada

Contents *continued*

Drug failure in reducing pressor effect of isometric handgrip stress test in hypertension 211

Sejan Lomid MD and Frederick W. Wolf MD Washington D C

The role of beta adrenoceptors in the coronary and systemic hemodynamic responses to emotional stress in conscious dogs 216

M Bergamasci PhD A M Caravaggi D V M V Mandelli D Sc and R G Shanks MD Milan Italy

Case reports

Cardiac lymphangioma and lipoma

Report of a case of simultaneous occurrence in association with lipomatous infiltration of the myocardium and cardiac arrhythmia 227

Daniel T. Ince MD and Gerald Fine MD Detroit Mich

Propranolol induced alopecia 236

Carrell M. Martin MD Major MC USA Edward G. Southwick MD Major MC USA and Howard I. Weibach MD San Francisco Calif

Clinical pathologic conference

Clinical pathologic conference 238

Yake A. Karachorlu MD Rolf M. Gunnar MD and Cecil A. Krahower MD Chicago Ill

Fundamentals of clinical cardiology

Mechanisms of cardiac arrhythmias From hypothesis to physiologic fact 249

Alfred Fick MD Chicago Ill

Appraisal and reappraisal of cardiac therapy

Drug therapy of heart disease in pediatric patients II

The treatment of congestive heart failure in infants and children with digitalis preparations 270

Monika M. Rudkowski MD Sanford V. Cohen MD and Eugene F. Doyle MD New York N Y

continued on page 7

V. L. 26 No. 2 August 1973 The American Heart Journal is published monthly by The C. V. Mosby Company
11830 Westlawn Industrial Drive, St. Louis, Mo 63141
A Circulation Periodical

| | U.S. | Canada \$ 10 | Other \$ 15 |
|----------------------------|---------|-----------------|----------------|
| Institutional (non-profit) | \$26.00 | \$ 9.50 | \$30.00 |
| Personal | \$19.50 | \$22.00 | \$ 3.25 |
| Student (reduced rate) | \$13.60 | \$16.65 | \$17.40 |

Second class postage paid at St. Louis, Mo.
Postmaster: Send address changes to The American Heart Journal, c/o The C. V. Mosby Company, 11830 Westlawn Industrial Drive, St. Louis, Mo 63141.

Subscription rates for institutions, libraries, schools, hospitals, and clinics only. For state, provincial, and national government bureaus and departments, and all commercial and private institutions and organizations, a separate rate schedule is available. For individual subscriptions, see the inside back cover of this journal.

Subscriptions may be ordered at any time.

Second class postage paid at St. Louis, Mo.

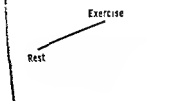
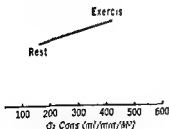
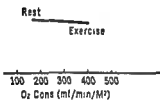
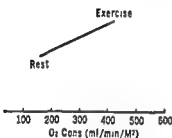
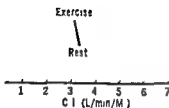
Printed in the U. S. A. Copyright © 1973 by The C. V. Mosby Company

MODIFYING

AND EXERCISING, BEFORE AND AFTER ISORDIL SUBLINGUAL

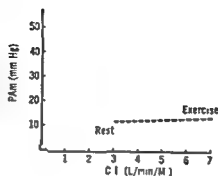
(isosorbide dinitrate)

BEFORE MEDICATION

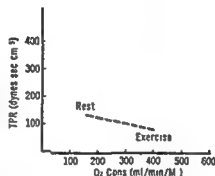


AFTER 5 mg ISORDIL SUBLINGUAL (isosorbide dinitrate)

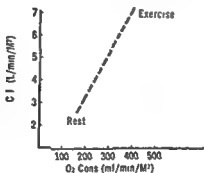
PULMONARY ARTERY (MEAN PRESSURE)



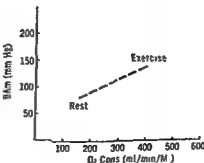
TOTAL PULMONARY RESISTANCE



CARDIAC INDEX



BRACHIAL ARTERY (MEAN PRESSURE)



TOTAL SYSTEMIC RESISTANCE



IN ANGINA PECTORIS

- to abort or terminate acute anginal attacks
- to reduce the frequency and severity of anginal episodes often caused by unavoidable everyday stress

ISORDIL SUBLINGUAL (ISOSORBIDE DINITRATE)

SUBLINGUAL TABLETS 25 mg and 5 mg

*Indications Based on a review of this drug by the National Academy of Sciences—National Research Council and/or other information FDA has classified the indication as follows

Probably effective When taken by the sublingual route Isordil Sublingual is indicated for the treatment of acute anginal attacks and for prophylaxis in situations likely to provoke such attacks Final classification of the less than effective indications requires further investigation

Contraindication Idiosyncrasy to this drug

Warnings Data supporting the use of nitrates during the early days of the acute phase of myocardial infarction (the period during which clinical and laboratory findings are unstable) are insufficient to establish safety

Precautions Tolerance to this drug and cross tolerance to other nitrites and nitrates may occur

Adverse Reactions Cutaneous vasodilation with flushing Headache is common and may be severe and persistent Transient episodes of dizziness and weakness as well as other signs of cerebral ischemia associated with postural hypotension may occasionally develop This drug can act as a physiological antagonist to norepinephrine acetylcholine histamine and many other agents An occasional individual exhibits marked sensitivity to the hypotensive effects of nitrite and severe responses (nausea vomiting weakness restlessness pallor perspiration and collapse) can occur even with the usual therapeutic dose Alcohol may enhance this effect Drug rash and/or exfoliative dermatitis may occasionally occur

Consult direction circular before prescribing

May we send you reprints detailed information and/or professional samples?

IVES LABORATORIES INC

N. York, N.Y. 10017

DEDICATED TO IMPROVING THE QUALITY OF LIFE THROUGH MEDICINE



Contents *continued*

Annotations

Ischemic cardiomyopathy 276

G E Burch MD New Orleans La

The nitroblue tetrazolium dye test and infection
in the renal patient 277

*Michael R Hollman MD and Denis R Miller MD
New York N Y*

The magnitude of risk of developing complete heart block
in patients with LAD RBBB 278

Henri E Kulbertus MD Liège Belgium

Renal excretion of sulfadimidine in normal and
uremic subjects 280

Erna Fischer MD Hille pd Denmark

Letters to the Editor

Cardioversion after valve replacement 282

Arthur Selzer MD San Francisco Calif

Reply 282

Hans Jenzer MD and Bernard Lown MD Boston Mass

✓ The Yogic claim of voluntary control over the heart
beat an unusual demonstration 282

*L A Kathari M Sc MA US Arun Borda MD and
O P Gupta MD Udaipur India*

Atropine in acute myocardial infarction 284

David Scheff MD New York N Y

Practolol in treating tachyarrhythmias 284

*Jean Pierre Van Durme MD Leo Bossaert MD Paul Vermeire MD
and René Pannier MD Gent Belgium*

Right atrial electrocardiogram resulting in hemopericardium 285

*Robert Hataf MD Claude Seban MD Richa d Benaim MD
and Paul Chiche MD FACC Paris France*

Atrial ectopic tachycardia 285

*W J Mandel MD J Lozano MD and H Hayakawa MD
Los Angeles Calif Caracas Venezuela and Tokyo Japan*

Reply 286

Bruce N Goldsayer MD Philadelphia Pa

Systolic time intervals in man 286

Jean M Pouget MD and Willis d S Harris MD Chicago Ill

Reply 286

*Valdemar Lindquist MB MRCP Richard D Spangler MD FCCP
and S Gilbert Blount Jr MD Denver Colo*

Book reviews

Book reviews 287

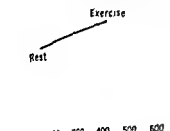
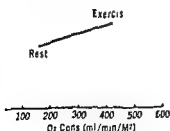
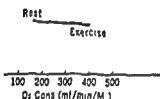
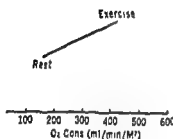
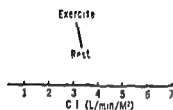
Books received

Books received 288

August

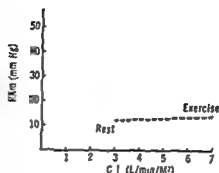
ND EXERCISING, BEFORE AND AFTER ISORDIL® SUBLINGUAL (isosorbide dinitrate)

BEFORE MEDICATION

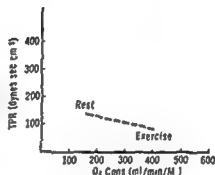


AFTER 5 mg ISORDIL SUBLINGUAL (isosorbide dinitrate)

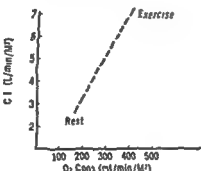
PULMONARY ARTERY (MEAN PRESSURE)



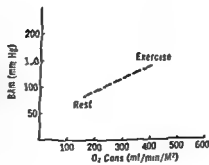
TOTAL PULMONARY RESISTANCE



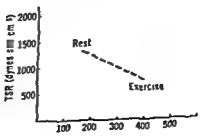
CARDIAC INDEX



BRACHIAL ARTERY (MEAN PRESSURE)



TOTAL SYSTEMIC RESISTANCE



IN ANGINA PECTORIS

- to abort or terminate acute anginal attacks
- to reduce the frequency and severity of anginal episodes often caused by unavoidable everyday stress

ISORDIL® SUBLINGUAL (ISOSORBIDE DINITRATE) SUBLINGUAL TABLETS 25 mg and 5 mg

*Indications Based on a review of this drug by the National Academy of Sciences—National Research Council and/or other information FDA has classified the indication as follows

Probably effective When taken by the sublingual route Isordil Sublingual is indicated for the treatment of acute anginal attacks and for prophylaxis in situations likely to provoke such attacks
Final classification of the less than effective indications requires further investigation

Contraindication Idiosyncrasy to this drug

Warnings Data supporting the use of nitrites during the early days of the acute phase of myocardial infarction (the period during which clinical and laboratory findings are unstable) are insufficient to establish safety

Precautions Tolerance to this drug and cross tolerance to other nitrites and nitrates may occur

Adverse Reactions Cutaneous vasodilation with flushing Headache is common and may be severe and persistent Transient episodes of dizziness and weakness as well as other signs of cerebral ischemia associated with postural hypotension may occasionally develop This drug can act as a physiological antagonist to norepinephrine acetylcholine histamine and many other agents An occasional individual exhibits marked sensitivity to the hypotensive effects of nitrite and severe responses (nausea vomiting weakness restlessness pallor perspiration and collapse) can occur even with the usual therapeutic dose Alcohol may enhance this effect Drug rash and/or exfoliative dermatitis may occasionally occur

Consult direction circular before prescribing

May we send you reprints detailed information and/or professional samples?

IVES LABORATORIES INC
New York, N.Y. 10017
DEDICATED TO IMPROVING THE QUALITY OF LIFE THROUGH MEDICINE



Adapted from Katoar
atory Evaluati
ic Exhibit

Pressure flow studies in man The nature of the aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome 350

W Eugene Kendall MD Judith C Rembert PhD and Joseph C Greenfield Jr MD Durham NC

Anodal stimulation as a cause of pacemaker induced ventricular fibrillation 366

Thomas A Preston MD Ann Arbor Mich

The value of warning arrhythmias in the prediction of ventricular fibrillation within one hour of coronary occlusion Experimental studies in the baboon 373

Koenraad J J Bruynel MD and Lionel H Opie MD Cape Town South Africa

Cardiovascular events in anxiety Experimental studies in the conscious dog 385

Mario Bergamaschi PhD and Anna M Longoni D Sc Milano Italy

Case reports

Cerebrovascular accident with unusual electrocardiographic changes 393

Gary J Anderson MD Robert Woodburn MD and Charles Fisch MD F.A.C.C Indianapolis Ind

Malfunction of a Cutter Smeloff mitral ball valve prosthesis Diagnosis by phonocardiography and echocardiography 399

Israel Belenkie MD Matthew Carr MD R C Schlant MD D O Nutter MD and P N Symbas MD Atlanta Ga

continued on page 7

V 186 N 3 Sept mbr 1973 Th American Heart Journal is published monthly by The C. V. Mosby Company
5130 W. 11th Industrial Dr. St. Louis, Mo 63141
A useful subscription list on p. 4

| | G S | Cash | Other |
|--------------------|---------|---------|---------|
| | | | amount |
| Subscription | \$ 6.50 | \$79.50 | \$30.75 |
| Postage | \$19.50 | \$.50 | \$73.25 |
| Subscription agent | \$13.65 | \$16.6 | \$17.40 |

1. If you are a subscriber, please send your money to the office of the publisher, The C. V. Mosby Company, 5130 W. 11th Industrial Dr., St. Louis, Mo 63141.

In addition to (multiple) subscribers, we are available to public and private libraries, schools, hospitals and all city, county, state, provincial and national government departments and all commercial and private institutions and organizations.

Personal subscription rates and rates for institutions are available on request in the name of and by the publisher.

Subscriptions may begin at any time.

Second class postage paid at St. Louis, Mo.

Printed in the U.S.A. Copyright © 1973 by The C. V. Mosby Company

"The drug of choice for oral replacement of potassium is potassium chloride solution."

AMA Drug Evaluations 1971 First Edition
Chicago: American Medical Association p. 121

Kay Ciel[®] Elixir is potassium chloride... tastes good too!

COMPOSITION Each 15 cc. (one tablespoonful) contains potassium chloride 15 Gm. supplying 80 meq. of elemental potassium in a cherry flavored palatable base alcohol 4% Contains no sugar

INDICATIONS Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy digitalis intoxication low dietary intake of potassium or as a result of excessive vomiting and diarrhea

CONTRAINDICATIONS Impaired renal function untreated Addison's Disease dehydration heat cramps and hyperkalemia

PRECAUTIONS Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient since the amount of deficiency and corresponding daily dose is

often not known. Excessive or even therapeutic doses may result in potassium intoxication. Patients should be frequently checked and periodic ECG and/or plasma potassium levels made. High plasma concentrations of potassium ion may cause cardiac depression, arrhythmias or arrest. Use with caution in patients with cardiac disease. In hypokalemic states, attention should be directed toward the correction of the frequently associated hypochloremic azotemia.

SIDE EFFECTS Vomiting, nausea, abdominal discomfort and diarrhea may occur. Symptoms and signs of potassium intoxication include listlessness, mental confusion, paresthesia of the extremities, weakness of the legs, flaccid paralysis, fall in blood pressure, cardiac arrhythmias and heart block. When hyperkalemia

exists, it should be promptly treated with the discontinuance of potassium administration or other step to lower serum levels. If indicated, since sudden shift in plasma levels may induce potentially dangerous cardiac arrhythmias.

DOSE AND ADMINISTRATION Adults one tablespoonful (15 cc.) diluted in one glass of water twice daily after the morning and evening meal. Larger doses may be indicated according to the individual patient's requirements but should be administered under close supervision due to the possibility of potassium intoxication. Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal injury.

HOW SUPPLIED One pint and one gallon bottles

Cooper

Cooper Laboratories Inc. Wayne, N.J. 07470/St. Thérèse, P.Q., Canada

Contents *continued*

Review

The use of spironolactone in the diagnosis and the treatment of hypertension associated with mineralocorticoid excess 404

D C Beeters MRCP J J Brown FRCP

*J B Ferriss MRCP R Fraser PhD A F Lister FRCP
and J I S Robertson Glasgow Scotland*

Fundamentals of clinical cardiology

Peripheral arterial occlusion in patients with acute coronary heart disease 415

*Sandor A Friedman MD FACP Mohendra Pandya MD and
Ermst Greif MD FACP Brooklyn N Y*

Appraisal and reappraisal of cardiac therapy

Indications for aortocoronary artery bypass surgery 420

Alfred J Hallman MD New York N Y

Annotations

Study of man himself 425

G E Burch MD New Orleans La

Longevity of athletes 425

George A Sheehan MD Red Bank N J

Dopamine test for the diagnosis of coronary insufficiency 426

*O Visio MD F N Eggerdy MD and G Malagnino MD
Parma Italy*

Complications of transfemoral coronary arteriography and their prevention using heparin 428

Kenneth M Fyer MD Seattle Wash

Letters to the Editor

Pacemaker catheter displacement 429

Isidore Allen MD New York N Y

Reply 429

Thomas A Preston MD Seattle Wash

Book reviews

Book reviews 430

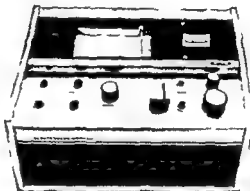
Announcements

Announcements 430

• Physician • Nurse • Biomedical Engineer • Administrator • Medical Technician



Each gets what
he wants—
in a new
Burdick EK/5A
electro-
cardiograph



THE PHYSICIAN
demands diagnostic accuracy, definition, dependability, no time consuming re takes, mistake free lead coding — and that's what the solid state EK/5A delivers. Exceeds AHA frequency response recommendations.

THE NURSE OR MEDICAL TECHNICIAN
wants the ease of operation that results in time saving—from features like the EK/5A automatic lead marking activated by a fastswitching lead selector and easy portability.

THE BIOMED ENGINEER
looks for multiple use in the hospital, patient isolated circuitry (leakage is less than 10 micro amperes), solid state simplicity, multiple circuit boards for easier, faster service, computer compatibility, sound ECG design.

THE ADMINISTRATOR AND PURCHASING AGENT
like (1) the savings that result from dependable performance (2) medical staff satisfaction and (3) Burdick dealer service and maintenance availability.
THE BURDICK CORPORATION
Milton, Wis 53563

BURDICK

Review

The use of spironolactone in the diagnosis and the treatment of hypertension associated with mineralocorticoid excess 401

D G Beevers MRCP J J Brown FRCP

J B Ferriss MRCP R Fraser PhD A F Lever FRCP
and *J I S Robertson Glasgow Scotland*

Fundamentals of clinical cardiology

Peripheral arterial occlusion in patients with acute coronary heart disease 415

Sandor I Friedman MD FACP Mahendra Pandya MD and
Ernst Grief MD FACP Brooklyn N Y

Appraisal and reappraisal of cardiac therapy

Indications for aortocoronary artery bypass surgery 420

Alfred J Kaltman MD New York N Y

Annotations

Study of man himself 425

G I Burch MD New Orleans La

Longevity of athletes 425

George A Sheehan MD Red Bank N J

Dopamine test for the diagnosis of coronary insufficiency 426

U Visiofi MD F N Effendi MD and G Malagnino MD
Parma Italy

Complications of transfemoral coronary arteriography and their prevention using heparin 428

Kenneth M Eyer MD Seattle Wash

Letters to the Editor

Pacemaker catheter displacement 429

Ludwig Klein MD New York N Y

Reply 429

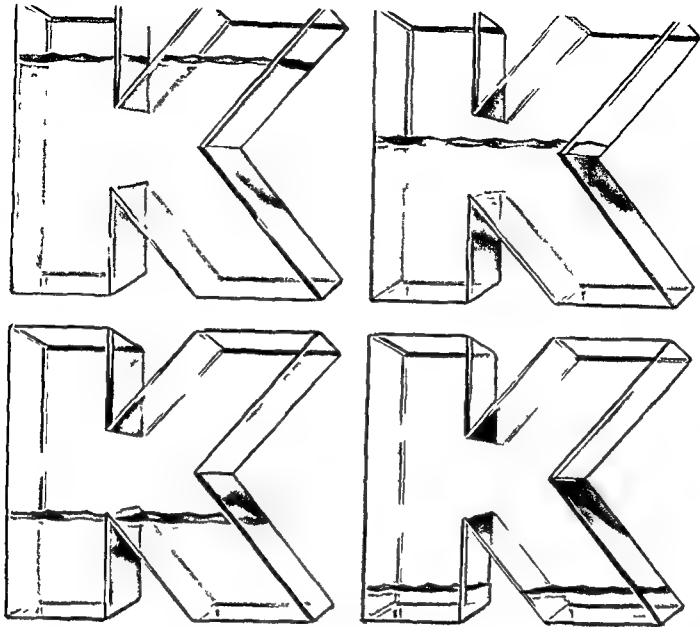
Thomas A Preston MD Seattle Wash

Book reviews

Book reviews 430

Announcements

Announcements 430



'Dyrenium' Conserves Potassium add it to the thiazides or furosemide

In the treatment of edema*

Dyrenium conserves the potassium most other diuretics waste and augments the action of other diuretics

Before prescribing see complete prescribing information in SK&F literature or PDR

Indications Edema associated with congestive heart failure cirrhosis and nephrotic syndrome steroid-induced edema idiopathic edema edema due to secondary hyperaldosteronism and edema resistant to other diuretic therapy

Contraindications Severe or progressive kidney disease or dysfunction (possible exception nephrosis) Severe hepatic disease Pre-existing elevated serum potassium Hypersensitivity to the drug Continued use in developing hyperkalemia Do not give potassium supplements either by drug or by diet

Warnings Observe regularly for possible blood dyscrasias liver damage or other

idiosyncratic reactions Blood dyscrasias have been reported Check BUN and serum potassium periodically especially in the elderly diabetics and those with suspected or confirmed renal insufficiency Use in pregnancy only when essential to patient welfare

Dyrenium (triamterene SK&F) and spironolactone are not usually used concurrently if they are however frequent serum potassium determinations are required

Precautions If hyperkalemia develops withdraw the drug The following may also occur electrolyte imbalance low salt syndrome (with low salt intake) reversible mild nitrogen retention decreasing alkali reserve with possible metabolic acidosis Do periodic hematologic studies in cirrhotics with splenomegaly Concomitant use with anti-hypertensive drugs may result in an additive hypotensive effect When Dyrenium is to be discontinued after intensive or prolonged therapy with draw gradually because of possible rebound kaliuresis

Adverse Reactions Diarrhea nausea and vomiting (may indicate electrolyte imbalance) other gastrointestinal disturbances weakness headache dry mouth anaphylaxis photosensitivity elevated uric acid rash

Note When combined with another diuretic the initial dosage of each agent should be lower than recommended
Supplied 100 mg capsules in bottles of 100

SK&F Co. Carolina PR 00630
A subsidiary of
Smith Kline & French Laboratories

100 mg capsules

Dyrenium
brand of triamterene

Contents

Editorial

Central venous pressure: Physiological significance
and clinical implications 431

Arthur C Guyton MD and Carl E Jones PhD
Jackson Miss

Clinical communications

Improvement of left ventricular asynergy following
aortocoronary bypass surgery related to preoperative
electrocardiogram and vectorcardiogram 438

Kurian Jacob MD Michel Chabot MD Jacques Saliot MD
and Lucien Campeau MD Montreal Canada

Wenckebach periods with repetitive block.
Evaluation with His bundle recording 444

Ramesh C Dhirga MD Kenneth M Rosen MD FACC
and Shahbudin H Rahimtoola FRCP Chicago Ill

Electrocardiographic findings in single ventricle
and related conditions 449

M Quera-Jiménez MD M Casanova Gomez MD C Castro Gutierrez MD
F Moreno-Granado MD V Pérez-Martínez MD and
G Merino-Baltes MD Madrid Spain

Radiological patterns of obstructive cardiomyopathy
of the left ventricle in childhood 462

C Pernet MD J C Hoefel MD M Henry MD
A M Worms MD and G Rothhahn MD
Dommarin Les Toul France

Embolic coronary artery occlusion in percutaneous
transfemoral coronary arteriography 467

Angel de la Torre MD Daniel Jacobs MD Juan Aleman MD
and George A Anderson MD Jacksonville Fla

Asymptomatic electrocardiographic alterations
in sarcoidosis 474

E. Sein MD I Jackler MD B Stimmel MD W Stern MD
and L E Siltbach MD New York NY

**"The drug of choice
for oral replacement
of potassium is
potassium chloride
solution."**

AMA Drug Evaluation 1971 First Edition
Chicago: American Medical Association p. 121

**Kay Ciel[®] Elixir is
potassium chloride...
tastes good too!**

COMPOSITION: Each 15 cc. (one table spoonful) contains potassium chloride 1.8 Gm. supplying 20 mEq. of elemental potassium in a cherry flavored palatable base. Alcohol 4%. Contains no sugar.

INDICATIONS: Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy; digitalis intoxication; low dietary intake of potassium or as a result of excessive vomiting and diarrhea.

CONTRAINDICATIONS: Impaired renal function; untreated Addison's Disease; dehydration; heat cramps; and hyperkalemia.

PRECAUTIONS: Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient since the amount of deficiency and corresponding daily dose is

often not known. Excessive or even therapeutic dosages may result in potassium intoxication. Patients should be frequently checked and periodic ECG and/or plasma potassium levels recorded. High plasma concentrations of potassium ion may cause cardiac depression, a rhythmless arrest. Use with caution in patients with cardiac disease. In hypokalemic states attention should be directed toward the correction of the frequently associated hypochloremic alkalosis.

SIDE EFFECTS: Vomiting, nausea, abdominal discomfort and diarrhea may occur. Symptoms and signs of potassium intoxication include listlessness, mental confusion, paresthesia of the extremities, weakness of the legs, flaccid paralysis, tachycardia, cardiac arrhythmias and heart block. When hyperkalemia

exists, it should be promptly treated with the discontinuance of potassium administration or other steps to lower serum levels if indicated. Since sudden shift in plasma levels may induce potentially dangerous cardiac arrhythmias.

DOSEAGE AND ADMINISTRATION: Adults: one tablespoonful (15 cc.) diluted in one glass of water, twice daily after the morning and evening meal. Larger dose may be indicated according to the individual patient's requirements but should be administered under close supervision due to the possibility of potassium intoxication. Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal injury.

HOW SUPPLIED: One pint and one gallon bottles.

Cooper

Cooper Laboratories Inc. Wayne, N.J. 07470/Sie Theresé, P.Q. Canada

Adverse reactions to propranolol in hospitalized medical patients. A report from the Boston Collaborative Drug Surveillance Program 478

*David J Greenblatt MD and Jan Koch-Weser MD
Boston Mass*

Experimental and laboratory reports

The relationship of coronary collateral inlet flow and retrograde flow in mongrel dogs 485

*Anthony A Cibulski MD BEE Patrick H Lehan MD
and Harper K Hellemis MD Jackson Miss*

Determination of systolic intervals utilizing the carotid first derivative 495

*Priya S Nandi MD and David H Spodick MD
Boston Mass*

Transient ST elevation detected by 24 hour ECG monitoring during normal daily activity 501

*Basil Golding MB ChB Eliana Wolf MD Dan Tzivoni MD
and Shlomo Stern MD Jerusalem Israel*

Depression of cardiac performance by ethanol unmasked during autonomic blockade 508

Maylene Wong MD Los Angeles Calif

Investigation of atrial aberration as a cause of altered P wave contour 516

*Peter Probst MD Jane Hunter AB Olive Gamble AB
and Keith Cohn MD San Francisco Calif*

continued on page 7

Vol 86 No 4 October 1973 The American Heart Journal is published monthly by The C. V. Mosby Co. 11830 W. Jefferson Blvd., Suite 100, St. Louis, MO 63141
Annual subscription rate

| | U.S. | Canada | Other countries |
|---|---------|---------|-----------------|
| Institutional | \$27.50 | \$30.50 | \$31.25 |
| Personal | \$19.50 | \$22.50 | \$23.25 |
| Third class postage paid at St. Louis, MO | \$13.65 | \$16.65 | \$17.40 |

Subscription price \$35.00 postpaid. Remittances should be made by check, draft, post office order, or money order payable to this journal.

Individual (multiple copies) subscriptions are available to public and private libraries, schools, hospitals, city, county, state, provincial, and national government bureaus and departments and all commercial and private institutions and organizations.

Personal subscriptions and all student rate subscriptions must be in the name of a subscriber with duals.

Subscriptions may be ordered at any time.

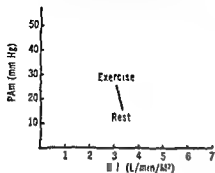
Second class postage paid at St. Louis, Mo.

Printed in the U.S.A. Copyright © 1973 by The C. V. Mosby Co.

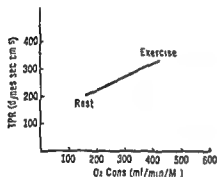
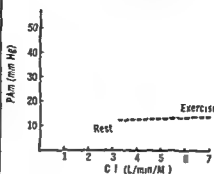
AND EXERCISING, BEFORE AND AFTER ISORDIL[®] SUBLINGUAL (isosorbide dinitrate)

BEFORE MEDICATION

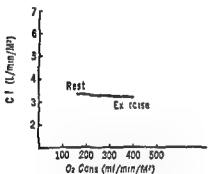
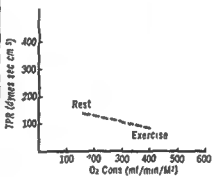
AFTER 5 mg ISORDIL SUBLINGUAL (isosorbide dinitrate)



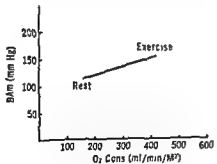
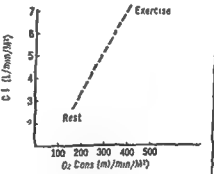
PULMONARY
ARTERY
(MEAN
PRESSURE)



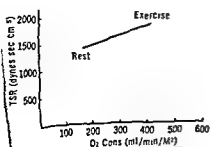
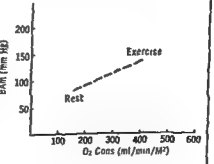
TOTAL
PULMONARY
RESISTANCE



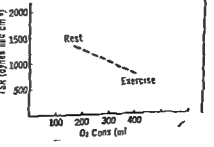
CARDIAC
INDEX



BRACHIAL
ARTERY
(MEAN
PRESSURE)



TOTAL
SYSTEMIC
RESISTANCE



IN ANGINA

- to abort or terminate acute anginal attacks
- to reduce the frequency and severity of anginal episodes often caused unavoidable everyday stress

ISORDIL[®] SUBLINGUAL

(ISOSORBIDE DINITRATE)
SUBLINGUAL TABLETS 2.5 mg and 5 mg

***Indications** Based on a review of this by the National Academy of Pharmaceutical Research Council and/or other information FDA has classified the indication follows

Probably effective When taken by sublingual route Isordil Sublingual is indicated for the treatment of acute attacks and for prophylaxis in situations likely to provoke such attacks Final classification of the less than effect indications requires further investigation

Contraindication Idiosyncrasy to this drug
Warnings Data supporting the use of during the early days of the acute phase myocardial infarction the period during which clinical and laboratory findings are usually insufficient to establish safety

Precautions Tolerance to this drug and tolerance to other nitrites and nitrates occur

Adverse Reactions Cutaneous vasodilation flushing Headache is common and may be severe and persistent Transient episodes of dizziness and weakness as well as other effects of cerebral ischemia associated with post hypotension may occasionally develop drug can act as a physiological antagonist norepinephrine acetylcholine histamine many other agents An occasional individual exhibits marked sensitivity to the hypotensive effects of nitrite and severe responses (sea vomiting weakness restlessness perspiration and collapse) can occur even the usual therapeutic dose Alcohol may enhance this effect Drug rash and/or exfoliative dermatitis may occasionally occur

Consult direction circular before prescribing
May we send you reprints detailed information and/or professional samples?

IVES LABORATORIES INC

New York, N.Y. 10017
DEDICATED TO IMPROVING THE QUALITY OF LIFE THROUGH MEDICINE

Adapted from
Laboratory Experiments
Scientific
June

Case reports

Viral infection of the aorta of man associated
with early atherosclerotic changes 523

*G E Burch MD J M Harb PhD Y Hiramoto MD
and Lana Shewey MD New Orleans La*

Charcot Marie Tooth disease Wolff Parkinson White
syndrome and abnormal intracardiac conduction 535

Dorance Bowers MD Kelowna B C Canada

Clinical pathologic conference

Clinical pathologic conference 539

*V Simon MD H Cohen MD G Gluck MD A Kanter MD
B Levin MD and C L Pirani MD Chicago Ill*

Fundamentals of clinical cardiology

The spectrum of common atrioventricular orifice (canal) 553

Saroja Bharati MD and Maurice Lev MD Chicago Ill

Appraisal and reappraisal of cardiac therapy

Drug therapy of heart disease in pediatric patients III

The therapeutic challenge of supraventricular
tachyarrhythmias in infants and children 562

*Monika M Rutkowski MD Eugene F Doyle MD
and Sanford N Cohen MD New York N Y*

Announcements

Theoretic considerations of the post systolic
dip of constrictive pericarditis 569

G E Burch MD and T D Giles MD New Orleans La

Blood flow in the internal mammary artery 570

Hendrick B Barner MD St Louis Mo

Cell growth in man 571

C G D Brook MD MRCP London England

Serum enzyme levels after operation 573

*Sylvia M Watkins BM MRCP and Adam Lewis FRCS
London England*

Book reviews

Book reviews 575

Announcements

Announcements 576

Patients at risk

When the patient is at risk because of K-loss

Potassium depletion can become a hazard in

- diuretic therapy
- digitalis therapy
- prolonged and severe vomiting
- prolonged corticosteroid therapy

an answer that makes sense is K-LOR
(POTASSIUM CHLORIDE SUPPLEMENT)

K-LOR provides both potassium and chloride ions allowing for complete correction of potassium imbalance associated with diuretic therapy

There is less chance of gastric upset with K-LOR since it is provided in single dose packets of powder that must be diluted with liquid

The single-dose packet assures you of accurate dosage and provides convenience for the patient

K-LOR™ (POTASSIUM CHLORIDE SUPPLEMENT)

Because potassium supplementation is serious medicine

Contraindications Potassium chloride is contraindicated in the presence of severe renal impairment with oliguria or azotemia; untreated Addison's disease; adynamia episodica hereditaria; acute dehydration; heat cramps; and hyperkalemia from any cause.

Potassium chloride should not be employed in patients receiving potassium sparing agents such as aldosterone antagonists and triamterene.

Precautions With normal kidney function, potassium intoxication from oral administration is not likely to occur, since renal excretion of the ion increases in response to a rise in the concentration of body potassium. Nevertheless, potassium supplements must be administered with caution, since the dietary or daily amount is not accurately known. Frequent checks of the patient's clinical status and periodic ECG and/or serum potassium levels should be done. High serum concentrations of potassium ion may result in death through cardiac depression, arrhythmia, or arrest. The drug should be used with caution in the presence of cardiac disease and systemic acidosis.

Adverse Reactions Side effects include abdominal discomfort, nausea, vomiting, and diarrhea.

In the presence of renal dysfunction, it may be possible to induce hyperkalemia by oral administration of potassium salts. The symptoms and signs of potassium intoxication include paresthesias of the extremities, weakness and heaviness of the legs, flaccid paralysis, little or no mental confusion, fall in blood pressure, cardiac arrhythmias and heart block. Electrocardiographic abnormalities such as disappearance of the P wave, widening and slurring of the QRS complex, changes of the ST segment and tall peaked T waves may be noted with hyperkalemia.

TM—T denotes

30405



K-LOR™ 20 meq
POTASSIUM CHLORIDE SUPPLEMENT
100 mg 40 mEq 20 meq 10 mEq 5 mEq
100 mg 40 mEq 20 meq 10 mEq 5 mEq
100 mg 40 mEq 20 meq 10 mEq 5 mEq

Contents

Editorial

Cyclophosphamide and the treatment of the
nephrotic syndrome in adults 577

C S Ogg M.D. MRCP and J S Cameron MD FRCP London England

Clinical communications

Early systolic notch in the apexcardiogram in mitral stenosis 582

*Lewis C Becker MD Andrew P Klaus MD and
J O Neal Humphries MD Baltimore Md*

An analysis of deaths occurring in association with
coronary arteriography 587

*Timothy Takaro MD Herbert N Hullgren MD David Littmann MD and
Elizabeth C Wright Olsen N C*

Left ventricular gallop sound and acute myocardial infarction 598

*Charles P Riley MD Richard O Russell Jr MD and
Charles E Rackley MD Birmingham Ala*

Coronary arteriographic findings in patients with axis shifts or
ST segment elevations on exercise stress testing 603

*Frederick N Hegge MD Nasip Tuna MD PhD and
Howard E Butchell MD PhD Minneapolis Minn*

Electrophysiological and histological abnormalities
of the heart in myotonic dystrophy 616

*Nobuhiro Uemura MD Hiromitsu Tanaka MD Tatsuru Aizawa MD
Aobuo Hashiguchi MD Masahiro Yoshimura MD Shinichi Terashi MD and
Takuya Kanehisa MD Kagoshima Japan*

Immunologic findings in idiopathic cardiomyopathy
A prospective serial study 625

*Allan B Kirsner MD Evelyn V Hett MD FACP and
Vivian O Fowler MD FACP Cincinnati Ohio*

Experimental and laboratory reports

The use of radio iodinated toluidine blue for myocardial scintigrams 631

*Edward A Carr Jr MD Mary Carroll BA Walter DiGiulio MD and
Donald C Blair MD Ann Arbor and Detroit Mich*

Patients at risk

When the patient is at risk because of K-loss

Potassium depletion can become a hazard in

- diuretic therapy
- digitalis therapy
- prolonged and severe vomiting
- prolonged corticosteroid therapy

an answer that makes sense is K-LOR
(POTASSIUM CHLORIDE SUPPLEMENT)

K-LOR provides both potassium and chloride ions
allowing for complete correction of potassium
imbalance associated with diuretic therapy

There is less chance of gastric upset with K-LOR since it is provided in single dose packets of powder that must be diluted with liquid

The single-dose packet assures you of accurate dosage and provides convenience for the patient

K-LOR™ (POTASSIUM CHLORIDE SUPPLEMENT)

Because potassium supplementation is serious medicine

Contraindications Potassium chloride is contraindicated in the presence of severe renal impairment with oliguria or azotemia untreated Addison's disease adynamia episodica hereditaria acute dehydration heat cramps and hyperkalemia from any cause

Potassium chloride should not be employed in patients receiving potassium sparing agents such as aldosterone antagonists and triamterene.

Precautions: With renal kidney function, potassium intoxication from oral administration is not likely to occur since renal excretion of the ion increases in response to a rise in the concentration of body potassium. Nevertheless potassium supplements must be administered with caution since the extra or daily amount is not accurately known. Frequent checks of the patient's clinical status and periodic ECG and/or serum potassium levels should be done. High serum potassium levels may result in death with caution in the presence of cardiac arrhythmia or arrest. The drug should be used with caution in the presence of cardiac disease and systemic acidosis.

Adverse Reactions Side effects include abdominal discomfort, nausea, vomiting, and diarrhea.

And in the presence of renal dysfunction it may be possible to induce hyperkalemia by oral administration of potassium salts. The symptoms and signs of potassium intoxication include paresthesias of the extremities weakness and heaviness of the legs flaccid paralysis listlessness mental confusion fall in blood pressure cardiac arrhythmias and heart block Electrocardiographic abnormalities such as disappearance of the P wave widening and slurring of the QRS complex changes of the S T segment and tall peaked T waves may be treated with hyperkalemia

TM=1 d m a k

Volume



Contents

Editorial

Cyclophosphamide and the treatment of the
nephrotic syndrome in adults 577

C S Orr MD MRCP and J S Cameron MD FRCP London England

Clinical communications

Early systolic notch in the apexcardiogram in mitral stenosis 582

*Lewis C Becker MD Andrew P Klaus MD and
J O Neal Humphries MD Baltimore Md*

An analysis of deaths occurring in association with
coronary arteriography 587

*Timothy Takaro MD Herbert N Hultgren MD David Luttmann MD and
Elizabeth C Wright Olsen NC*

Left ventricular gallop sound and acute myocardial infarction 598

*Charles P Raley MD Richard O Russell Jr MD and
Charles E Rackley MD Birmingham Ala*

Coronary arteriographic findings in patients with axis shifts or
S T segment elevations on exercise stress testing 603

*Frederick N Hegge MD Naoh Tuna MD PhD and
Howard B Burchell MD PhD Minneapolis Minn*

Electrophysiological and histological abnormalities
of the heart in myotonic dystrophy 616

*Nobuhiko Uemura MD Hiromitsu Tanaka MD Taisuru Numura MD
Nobuo Hashiguchi MD Masahiro Yoshimura MD Shinichi Terashi MD and
Takuya Kanehisa MD Kagoshima Japan*

Immunologic findings in idiopathic cardiomyopathy
A prospective serial study 625

*Allan B Kiersner MD Evelyn I Hess MD FACP and
Noble O Fowler MD FACP Cincinnati Ohio*

Experimental and laboratory reports

The use of radio iodinated toluidine blue for myocardial scintigrams 631

*Edward A Carr Jr MD Mary Carroll BA Walter DiGiulio MD and
Donald C Blair MD Ann Arbor and Detroit Mich*

**“The drug of choice
for oral replacement
of potassium is
potassium chloride
solution.”**

AMA Drug Evaluations 1971 First Edition
Chicago: American Medical Association p. 121

**Kay Ciel[®] Elixir is
potassium chloride...
tastes good too!**

COMPOSITION Each 15 cc (one table spoonful) contains potassium chloride 15 Gm. supplying 20 mEq of elemental potassium in a cherry flavored palatable base alcohol 4%. Contains no sugar.

INDICATIONS Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy digitalis intoxication low dietary intake of potassium or as a result of excessive vomiting and diarrhea.

CONTRAINDICATIONS Impaired renal function untreated Addison's Disease dehydration heat cramps and hyperkalemia.

PRECAUTIONS Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient since the amount of deficiency and corresponding daily dose is

often not known. Excessive or even therapeutic dosages may result in potassium intoxication. Patients should be frequently checked and periodic ECG and/or plasma potassium levels made. High plasma concentrations of potassium ion may cause cardiac depression, arrhythmias or arrest. Use with caution in patients with cardiac disease. In hypokalemic states attention should be directed toward the correction of the frequently associated hypochloremic alkalosis.

SIDE EFFECTS Vomiting, nausea, abdominal discomfort and diarrhea may occur. Symptoms and signs of potassium intoxication include listlessness, mental confusion, paresthesia of the extremities, weakness of the legs, flaccid paralysis, fall in blood pressure, cardiac arrhythmias and heart block. When hyperkalemia

exists, it should be promptly treated with the discontinuance of potassium administration or other steps to lower serum levels if indicated, since sudden shift in plasma levels may induce potentially dangerous cardiac arrhythmias.

DOSE AND ADMINISTRATION Adults one tablespoonful (15 cc) diluted in one glass of water, twice daily after the morning and evening meal. Larger doses may be indicated according to the individual patient's requirements but should be administered under close supervision due to the possibility of potassium intoxication. Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal injury.

HOW SUPPLIED One pint and one gallon bottles.

Cooper

Cooper Laboratories Inc. Wayne, N.J. 07470/Ste. Theresa P.Q. Canada

Disturbed blood flow in the carotid artery
Its physiological and clinical significance 644

Isaac Starr M D Christophe Ambrosi M D Joel H Manchester M D and
James C Shelburne M D Philadelphia Pa

Reciprocal movement of the right and left heart
demonstrated by directional Doppler ultrasound 651

Dennis Abelson M D and Hans R Muller M D Philadelphia Pa and Basel Switzerland

The effect of noradrenaline on blood flow and oxygen consumption
in normal and ischemic areas of myocardium 653

R. J Marshall B Sc and J R Parrott B Pharm M Sc Ph D Glasgow Scotland

The effect of heroin and multiple drug abuse on the electrocardiogram 663

Janet Lipski M D Barry Stimmel M D and
Ephraim Donoso M D New York N Y

Encephalomyocarditis (EMC) virus infection of the mouse aorta
An ultrastructural study 669

G E Burch M D and J M Horb Ph D New Orleans La

Case reports

Spontaneous return of sinus rhythm in older patients with
chronic atrial fibrillation and rheumatic mitral valve disease
Description of three patients 676

Thomas J Zimmerman M D Lofly L Bado M D U R C P and
Lewis F January M D Iowa City Iowa

Mitral disc variance (Harken prosthesis) 681

Sukh Dev Sharma M D Robert M Easley Jr M D Lawrence I Zaroff M D and
Sidney Goldstein M D Rochester N Y

continued on page 7

V. L. 80 No. 5 November 1973 The American Heart Journal is published monthly by The C. V. Mosby Company
11830 Weyburne Industrial Drive St. Louis, Mo. 63143
A special subscription price is available

| | <i>U.S.</i> | <i>Can. edn. Mex. Co.</i> | <i>Other countries</i> |
|----------------------------------|-------------|-------------------------------|----------------------------|
| Institutional | \$27.50 | \$30.50 | \$31.00 |
| Personal | \$19.50 | \$22.50 | \$23.25 |
| *Include 1 international postage | \$13.65 | \$16.65 | \$17.40 |

* Single copies are \$3.50 postpaid. Remittances should be made by check, draft or office or express money order payable to this Journal.

Institutional (multiple read) subscriptions are available to public and private libraries, schools, hospitals and clinics, city, county, state, provincial and national government, departments and all commercial and private institutions and organizations.

Personal subscription and all other subscriptions must be in the names of and billed to individuals.

* Subscriptions in foreign countries.

Second class postage paid at St. Louis, Mo.

Printed in the U. S. A. Copyright © 1973 by The C. V. Mosby Company

**"The drug of choice
for oral replacement
of potassium is
potassium chloride
solution."**

AMA Drug Evaluations 1971 First Edition
Chicago: American Medical Association ■ 121

**Kay Ciel® Elixir is
potassium chloride...
tastes good too!**

COMPOSITION Each 15 cc (one 1/2 bottle spoonful) contains potassium chloride 1.5 Gm. supplying 30 mEq of elemental potassium in a cherry flavored palatable base alcohol 4%. Contains no sugar.

INDICATIONS Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy digitalis intoxication low dietary intake of potassium or as a result of excessive vomiting and diarrhea.

CONTRAINDICATIONS Impaired renal function untreated Addison's Disease dehydration heat cramps and hyperkalemia.

PRECAUTIONS Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient since the amount of deficiency and corresponding daily dose is

often not known. Excessive or even therapeutic dosages may result in potassium intoxication. Patients should be frequently checked and periodic ECG and/or plasma potassium levels made. High plasma concentrations of potassium ion may cause cardiac depression arrhythmias or arrest. Use with caution in patients with cardiac disease. In hypokalemic states attention should be directed toward the correction of the frequently associated hypochloremic alkalosis.

SIDE EFFECTS Vomiting nausea abdominal discomfort and diarrhea may occur. Symptoms and signs of potassium intoxication include listlessness mental confusion paresthesia of the extremities weakness of the legs flaccid paralysis fall in blood pressure cardiac arrhythmias and heart block. When hyperkalemia

exists it should be promptly treated with the discontinuance of potassium administration or other steps to lower serum levels if indicated since sudden shifts in plasma levels may induce potentially dangerous cardiac arrhythmias.

DOSAGE AND ADMINISTRATION Adults one tablespoonful (15 cc.) diluted in one glass of water twice daily after the morning and evening meal. Larger dose may be indicated according to the individual patient's requirements but should be administered under close supervision due to the possibility of potassium intoxication. Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal injury.

HOW SUPPLIED One pint and one gallon bottles.

Cooper

Cooper Laboratories Inc. Wayne, N.J. 07470/St. Theresa P.Q. Canada

Review

- Nonpenetrating cardiac injuries A collective review 687
A James Liedtke MD and William E DeMuth Jr MD Hershey Pa

Fundamentals of clinical cardiology

- Echocardiographic manifestations of valvular vegetations 698
James C Dillon MD Harvey Feigenbaum MD Lee L Konecke MD
Richard H Davis MD and Sonia Chang AB Indianapolis Ind

Appraisal and reappraisal of cardiac therapy

- Complications of aortocoronary artery bypass surgery 705
Alfred J Kaitman MD New York NY

Annotations

- Whither electrocardiography? 709
J Scott Butterworth MD and Ephraim Glassman MD New York NY
- Aspirin in the prevention of thrombosis 711
J R O'Brien and R J H Butterfield Portsmouth and Nottingham England
- Left hemiblocks revisited from the histopathological viewpoint 712
J C Demoulin MD and H F Hulbertus MD Liege Belgium
- Fork and hypertension 713
George E Burch MD New Orleans La

Letters to the Editor

- ✓ Digoxin tablets A possible problem with biological availability 715
Assistant to the Director for Medical Communications
Food and Drug Administration Rockville Md

Book reviews

- Book reviews 716

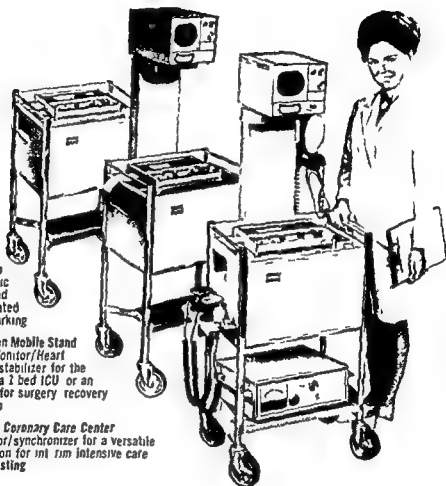
Books received

- Books received 717

Announcements

- Announcements 718

MONITORING VERSATILITY TAILORED TO YOUR NEEDS



Electrocardiograph
Start with the basic
mobile ECG — solid
state patient isolated
automatic lead marking

ECG with Monitor on Mobile Stand
Add the CS/515 Monitor/Heart
Rate Meter and a stabilizer for the
stand to give you a 1 bed ICU or an
ideal combination for surgery recovery
or emergency room

Complete Compact Coronary Care Center
Add the defibrillator/synchronizer for a versatile
low cost combination for int rim intensive care
or graded stress testing

BURDICK'S MOBILE COMPACT CORONARY CARE CENTER

Burdick has developed a sensible and economical approach to building your monitoring capabilities tailored to need. It answers the three demands of sound operational economy — **mobility, versatility, and equipment utilization** between departments. Using the proven concepts of the Burdick CCCC you **buy the components as you need them**.

For a brochure and additional supportive data contact your Burdick dealer or write The Burdick Corporation, Milton, Wis. 53563

BURDICK®

Contents

Acknowledgment to reviewers

Acknowledgment to reviewers 719

Editorial

An assessment of human cardiac transplantation 721

A Montem A Fadali MD and Louis A Soleff MD
Philadelphia Pa

Clinical communications

Electrocardiogram and vectorcardiogram in ventricular inversion (corrected transposition) 733

Benjamin E Victorica MD B Lynn Miller MD and
Ira H Gessner MD Gainesville Fla

Level of the base of the mitral valve 745

R W Brower PhD X H Krauss MD and G T Meester MD
Rotterdam The Netherlands

Non paroxysmal A V junctional tachycardia associated with acute myocardial infarction 754

Jaco Fishenfeld MD Kenneth B Desser MD and
Alberto Benchimol MD Phoenix Ari

Clinical diagnosis of persistent left superior vena cava by observation of jugular pulses 759

Simon Horvath MD Jose Esquivel A MD Fausto Altie MD
Eulo Lups H MD and Jorge Espino-Vela MD
Mexico City Mexico

Automatic ECG and blood pressure measurement in multitestung
Correlation of blood pressure and ECG abnormalities 764

H M Hochberg MD M F D George B SEE E L Schmalbach
and C A Caceres MD Cranbury N J

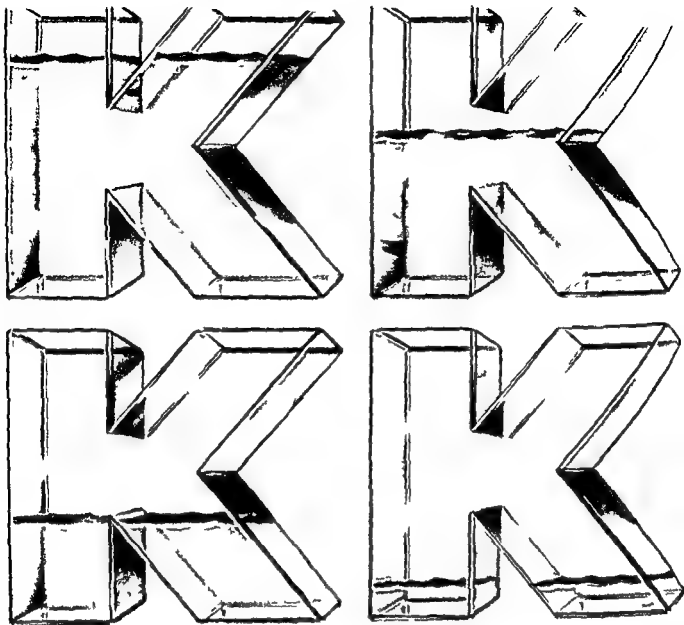
Experimental and laboratory reports

Electrophysiological evaluation of disopyramide in man 771

Mark E Josephson MD Anthony R Caracka MD Sun H Law MD
John J Gallagher MD and Anthony N Damato MD
Staten Island N Y

Are patients with essential hypertension and low renin
protected against stroke and heart attack? 781

Roland Stroobandt MD Robert Fagard MD and
Antoon A P C Amery MD Leuven Belgium



'Dyrenium' Conserves Potassium add it to the thiazides or furosemide

In the treatment of edema*

Dyrenium conserves the potassium most other diuretics waste and augments the action of other diuretics

Before prescribing see complete prescribing information in SK&F literature or PDR

Indications Edema associated with congestive heart failure, cirrhosis and nephrotic syndrome, steroid-induced edema, idiopathic edema, edema due to secondary hyperaldosteronism and edema resistant to other diuretic therapy

Contraindications Severe or progressive kidney disease or dysfunction (possible exception: nephrosis). Severe hepatic disease. Pre-existing elevated serum potassium. Hypersensitivity to the drug. Continued use in developing hyperkalemia. Do not give potassium supplements either by drug or by diet.

Warnings Observe regularly for possible blood dyscrasias, liver damage or other

idiosyncratic reactions. Blood dyscrasias have been reported. Check BUN and serum potassium periodically, especially in the elderly, diabetics and those with suspected or confirmed renal insufficiency. Use in pregnancy only when essential to patient welfare.

Dyrenium (triamterene, SK&F) and spironolactone are not usually used concurrently if they are, however, frequent serum potassium determinations are required.

Precautions If hyperkalemia develops, withdraw the drug. The following may also occur: electrolyte imbalance, low salt syndrome (with low salt intake), reversible mild nitrogen retention, decreasing alkali reserve with possible metabolic acidosis. Do periodic hematologic studies in cirrhotics with splenomegaly. Concomitant use with anti-hypertensive drugs may result in an additive hypotensive effect. When Dyrenium is to be discontinued after intensive or prolonged therapy, withdraw gradually because of possible rebound kaliuresis.

Adverse Reactions Diarrhea, nausea and vomiting (may indicate electrolyte imbalance), other gastrointestinal disturbances, weakness, headache, dry mouth, anaphylaxis, photosensitivity, elevated uric acid, rash.

Note When combined with another diuretic, the initial dosage of each agent should be lower than recommended.

Supplied 100 mg capsules in bottles of 100.

SK&F Co., Carolina, PR 00630
A subsidiary of
Smith, Kline & French Laboratories

100 mg capsules

Dyrenium®
brand of triamterene

Regional coronary flow with increased right ventricular output in anesthetized dogs 788

David E Fixler MD Joseph P Archie Jr MD
Daniel J Ulyet MD and Julius I F Hoffman MD
Dallas Texas and San Francisco Calif

Experimental myocardial infarction in the closed-chest dog
Controlled production of large or small areas of necrosis 798

Michael V Cohen MD and Per Eldh MD Boston Mass

Protection against epinephrine induced myocardial necrosis with clofibrate 805

Jacob I Haft MD Paul D Kran MD Frank Albert MD
and Rolf Oestrescher MD Bronx N Y

Case reports

Congenital absence of the circumflex coronary artery
Clinical and cinearteriographic observations 811

Vincent Barresi MD Armando Susmano MD Michael A Colandrea MD
Maurice L Bogdanoff MD and Joseph J Muenster MD
Chicago Ill

Traumatic coronary arterial fistula A case report
and review of the literature 817

Richard R Liberthson MD Kenneth Barron MD J Warren Harthorne MD
Robert E Dinsmore MD and Willard M Daggett MD
Boston Mass

Clinical pathologic conference

Clinical pathologic conference 822

Jane Somerville MD F.R.C.P S U Khalig M.R.C.Path
A C Brewer F.R.C.S and Donald Heath MD F.R.C.P F.R.C.Path
Liverpool England

continued on page 7

V. 1, No. 6 December 1973 The American Heart Journal is published monthly by The C. V. Mosby Company
11830 W. Line Industrial Drive St. Louis, Mo 63141
A. L. B. (p) 1183

| | U.S. | Canada Mexico | Other countries |
|----------------------|---------|------------------|--------------------|
| Institutional | \$27.50 | \$30.50 | \$31.25 |
| Personal | \$19.50 | \$22.50 | \$23.25 |
| Student (individual) | \$13.65 | \$16.65 | \$17.40 |

41 copies are \$3.50 postpaid. Payment may be made by check, draft, postal order or money order payable to the journal.

Institutional (multiple reader) subscriptions are available to public and private libraries, schools, hospitals, and libraries, city, county, state, provincial and national government bureaus and departments and all commercial and private institutions and organizations.

Personal subscription is a full student rate subscription and will be in the names of individuals.

Subscriptions may begin at any time.

Second class postage paid at St. Louis, Mo.

Printed in the U.S.A. Copyright © 1973 by The C. V. Mosby Company

"The drug of choice for oral replacement of potassium is potassium chloride solution."

AMA Drug Evaluations 1971 First Edition
Chicago American Medical Association p 121

Kay Ciel[®] Elixir is potassium chloride... tastes good too!

COMPOSITION Each 15 cc (one table spoonful) contains potassium chloride 1.5 Gm supplying 20 mEq of elemental potassium in a cherry flavored palatable base alcohol 4% Contains no sugar

INDICATIONS Treatment of potassium deficiency occurring especially during thiazide diuretic or corticosteroid therapy digitalis intoxication low dietary intake of potassium or as a result of excessive vomiting and diarrhea

CONTRAINDICATIONS Impaired renal function untreated Addison's Disease dehydration heat cramps and hyperkalemia

PRECAUTIONS Potassium chloride should be administered with caution and adjusted to the requirements of the individual patient since the amount of deficiency and corresponding daily dose is

often not known Excessive or even therapeutic dosages may result in potassium intoxication Patients should be frequently checked and periodic ECG and/or plasma potassium levels made High plasma concentrations of potassium ion may cause cardiac depression arrhythmias or arrest Use with caution in patients with cardiac disease in hypokalemic states attention should be directed toward the correction of the frequently associated hypochloremic alkalosis

SIDE EFFECTS Vomiting nausea abdominal discomfort and diarrhea may occur Symptoms and signs of renal intoxication include listlessness mental confusion paresthesia of the extremities weakness of the legs flaccid paralysis fall in blood pressure cardiac arrhythmias and heart block When hyperkalemia

exists it should be promptly treated with the discontinuance of potassium administration or other steps to lower serum levels if indicated since sudden shift in plasma levels may induce potentially dangerous cardiac arrhythmias

DOSAGE AND ADMINISTRATION Adults one tablespoonful (15 cc) diluted in one glass of water twice daily after the morning and evening meal Larger doses may be indicated according to the individual patient requirements but should be administered under close supervision due to the possibility of potassium intoxication Patients should be cautioned to follow directions explicitly in regard to dilution of Kay Ciel Elixir to prevent gastrointestinal injury

HOW SUPPLIED One pint and one gallon bottles



Cooper Laboratories Inc Wayne NJ 07470/Ste Therese PQ Canada

Contents *continued*

Fundamentals of clinical cardiology

Straight back syndrome Clinical and hemodynamic study of 9 cases 828

Shu o Matsuo MD Masao Yoshioka MD Katsutake Yano MD and Kunitake Hashiba MD Nagasaki Japan

Appraisal and reappraisal of cardiac therapy

Therapeutic uses of atrial pacing 835

Seymour Furman MD Bronx N Y

Annotations

Tricyclic antidepressants and cardiac disease 841

D C Moir MD Aberdeen Scotland

Hemoglobin plasma lipids and coronary heart disease 842

L E Bolliger MD Stockholm Sweden and L A Carlson MD Uppsala Sweden

Aspirin like drugs and prostaglandins 844

R J Flower London England

Surgical versus medical management of coronary heart disease 846

George E Burch MD New Orleans La

Letters to the Editor

Echocardiographic studies of mitral valve 847

Joram Glaser MD Cleveland Ohio

Reply 848

David A Mulward MD Lambert P McLaurin MD and Ernest Crange MD Chapel Hill N C

Heparin and venous thrombosis 849

J ■ Sharnoff MD Mount Vernon N Y

Reply 849

J R O'Brien MA DM Portsmouth England

Book reviews

Book reviews 851

Books received

Books received 851

Announcements

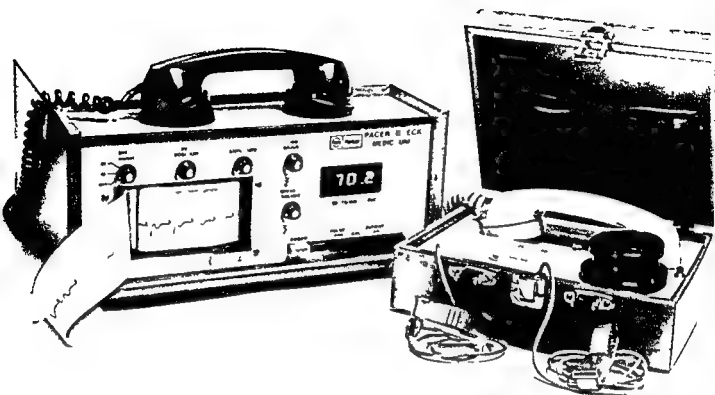
Announcements 852

Index

Author index 855

Subject index 861

December 1973



Make Life Easier For You And Your Pacer Patients

Now with the advent of the ESB Medcor Pacer-Check system it is possible for your office to conduct regular accurate recordings of heart pacer patient's ECG and pacer rates by telephone.

Both you and your patients benefit. Now with no more than a phone call you can have information which permits elective decisions on replacement rather than emergency or prophylactic replacements. (By using a regular patient checking program maximum pacer life can be realized.) Your patient and you have flexibility and safety at the same time.

This two component system operates simply requiring minimal training to both your personnel and patients.

The Medic Unit at left is the central component and is usually located in the hospital or clinic. It receives ECG and pacer rate input via unaltered telephones from the Patient Unit which is usually located in the patient's home. ECG data produced on a standard paper chart indicates the relationship between the QRS complex and

pacer artifact to ascertain pacer capture. Pacer rate is shown continuously by illuminated numbers on the front panel. This is the first system available to physicians that records ECG and simultaneously displays pacer rate.

The Patient Unit is lightweight and battery powered. It is capable of transmitting from any telephone to your office. Transmissions have been successfully made from such great distances as Tokyo to New York. To operate the patient places the telephone headset in the receiving cradle and then slips the two expandable bracelets over his forearms. Transmission begins immediately.

The Pacer Check system approved for reimbursement by Medicare has proven itself a useful and self supporting method of maintaining a desirable pacer program.

If the unique qualities of the Pacer Check system interest you contact ESB Medcor Inc. P.O. Box 6699 Hollywood Florida 33021 Tel: (305) 791 8310

- A unique new pacer electrode will be introduced at the American Heart Association meeting in Atlantic City, N.J.
- Visit us at booth C17-18 for a firsthand look

Editorial

Dietary treatment of renal failure

David S David M D *

New York N Y

In order for medicine to become a science one must attempt to quantitate. This is one of the many pearls of wisdom taught to me by the late Professor David Seegal in medical school. Although nephrology is far from being a true science there are many parameters which are readily measurable—i.e. weight intake and output, serum and urinary electrolytes, creatinine clearance, etc. Furthermore the nephrologist has at his disposal the ability to replace or replenish quantitatively any losses of fluids and electrolytes and with diuretics and dialysis the means to rid the body of any excesses of fluids and electrolytes. In other words the physician taking care of the patient with renal failure has the ability to regulate and take over through dietary means many of the kidney's normal functions. It is important for all physicians to master this skill because of the new hope that dialysis and transplantation have offered in prolonging and improving the quality of life of patients with renal failure.

Methods

Protein Since one of the main sources of uremic toxins contributing to the uremic syndrome is the protein in the diet, the

intake of protein has to be restricted. In patients with chronic renal disease protein restriction usually need not be imposed until the creatinine clearance falls below 10 to 15 cc per minute. Certain patients with renal disease who have associated catabolic states (postoperative infection and fever), excess non-dietary sources of protein (gastrointestinal bleeding) and low flow states (congestive heart failure, dehydration) may develop uremic symptoms at higher creatinine clearances and thereby will require protein restriction earlier. But it is important to note that patients with azotemia secondary to low flow states alone (without significant underlying renal disease) such as is commonly seen in congestive heart failure¹ rarely develop uremic symptoms requiring protein restriction and their azotemia responds very well to the therapy of the underlying disease.

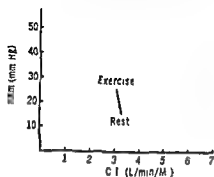
The Giovannetti diet and its modifications used in patients with renal failure is based on the observation that the high urea levels in the blood and in the colon are broken down by bacterial ureases in the colon to ammonia.¹ The ammonia is then taken up by 75% of the enterohepatic cir-

[illegible]

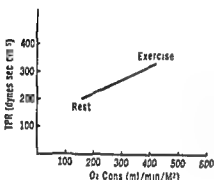
*Doc of CT law of psychology. Like floor advice to social 93le at Courtin's College of Physician

AND EXERCISING, BEFORE AND AFTER ISORDIL[®] SUBLINGUAL (isosorbide dinitrate)

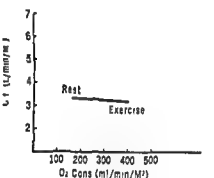
BEFORE MEDICATION



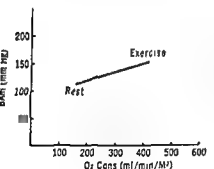
PULMONARY
ARTERY
(MEAN
PRESSURE)



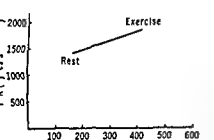
TOTAL
PULMONARY
RESISTANCE



CARDIAC
INDEX

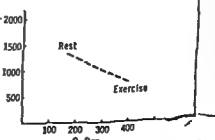
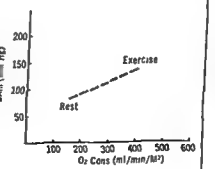
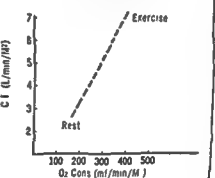
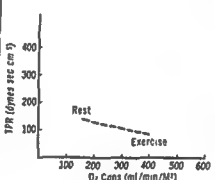
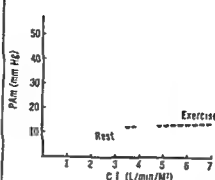


BRACHIAL
ARTERY
(MEAN
PRESSURE)



TOTAL
SYSTEMIC
RESISTANCE

AFTER 5 mg ISORDIL SUBLINGUAL (isosorbide dinitrate)



IN ANGINA PECTORIS

- to abort or terminate acute anginal attacks
- to reduce the frequency and severity of anginal episodes often caused by unavoidable everyday stress

ISORDIL[®] SUBLINGUAL

(ISOSORBIDE DINITRATE)
SUBLINGUAL TABLETS 2.5 mg and 5 mg

*Indications: Based on a review of this drug by the National Academy of Sciences—National Research Council and/or other information FDA has classified the indication as follows:

Probably effective: When taken by the sublingual route, Isordil Sublingual is indicated for the treatment of acute anginal attacks and for prophylaxis in situations likely to provoke such attacks. Final classification of the less than effective indications requires further investigation.

Contraindication: Idiosyncrasy to this drug.

Warnings: Data supporting the use of nitrites during the early days of the acute phase myocardial infarction (the period during which clinical and laboratory findings are unstable) are insufficient to establish safety.

Precautions: Tolerance to this drug and cross-tolerance to other nitrites and nitrates may occur.

Adverse Reactions: Cutaneous vasodilation, flushing, headache, are common and may be severe and persistent. Transient episodes of dizziness and weakness as well as other signs of cerebral ischemia associated with postural hypotension may occasionally develop. This drug can act as a physiological antagonist to norepinephrine, acetylcholine, histamine, and many other agents. An occasional individual exhibits marked sensitivity to the hypotensive effects of nitrite and severe responses (nausea, vomiting, weakness, restlessness, pallor, perspiration, and collapse) can occur even with the usual therapeutic dose. Alcohol may enhance this effect. Drug rash and/or exfoliative dermatitis may occasionally occur.

Consult direction circular before prescribing.

May we send you reprints, detailed information and/or professional samples?

IVES LABORATORIES INC.

New York, N.Y. 10017

DEDICATED TO IMPROVING THE QUALITY OF LIFE THROUGH MEDICINE



Adapted from
JAMA

Editorial

Dietary treatment of renal failure

David S David MD*
New York N Y

In order for medicine to become a science one must attempt to quantitate. This is one of the many pearls of wisdom taught to me by the late Professor David Seegal in medical school. Although nephrology is far from being a true science there are many parameters which are readily measurable—i.e. weight intake and output serum and urinary electrolytes creatinine clearance etc. Furthermore the nephrologist has at his disposal the ability to replace or replenish quantitatively any losses of fluids and electrolytes and with diuretics and dialysis the means to rid the body of any excesses of fluids and electrolytes. In other words the physician taking care of the patient with renal failure has the ability to regulate and take over through dietary means many of the kidney's normal functions. It is important for all physicians to master this skill because of the new hope that dialysis and transplantation have offered in prolonging and improving the quality of life of patients with renal failure.

Methods

Protein. Since one of the main sources of uremic toxins contributing to the uremic syndrome is the protein in the diet the

intake of protein has to be restricted. In patients with chronic renal disease protein restriction usually need not be imposed until the creatinine clearance falls below 10 to 15 cc per minute. Certain patients with renal disease who have associated catabolic states (postoperative infection and fever) excess non-dietary sources of protein (gastrointestinal bleeding) and low flow states (congestive heart failure dehydration) may develop uremic symptoms at higher creatinine clearances and thereby will require protein restriction earlier. But it is important to note that patients with azotemia secondary to low flow states alone (without significant underlying renal disease) such as is commonly seen in congestive heart failure¹ rarely develop uremic symptoms requiring protein restriction and their azotemia responds very well to the therapy of the underlying disease.

The Giovannetti diet and its modifications used in patients with renal failure is based on the observation that the high urea levels in the blood and in the colon are broken down by bacterial ureases in the colon to ammonia.^{2,3} The ammonia is then taken up by way of the enterohepatic cir-

From the Department of Medicine, Lusk Hospital Center, Columbia College of Physicians and Surgeons and the Renal Disease Treatment Center, New York Hospital-Cornell Medical Center.
Presented at the 1972 Annual Meeting of the American Medical Association.
Received for publication August 1, 1972.
Revised for publication October 1, 1972.
Direct of Clinical Nephrology, St. Luke Hospital Center, 1111 Lexington Avenue, New York, N.Y. 10017.
*Director of Clinical Nephrology, St. Luke Hospital Center, 1111 Lexington Avenue, New York, N.Y. 10017.

culation to the liver, which then synthesizes the non essential amino acids. The diet then need provide only the essential amino acids, and its protein content can be restricted to 0.3 to 0.5 Gm per kilogram of body weight.^{4,5} Egg chicken, beef steak and lamb, because of their high biological value, constitute the main proteins allowed in the diet.^{6,7}

The success of the Giovannetti diet, like any other diet depends on whether the patient adheres to it or not. There is one very important factor that the physician has under his control that will insure the success of the diet. This is to get a knowledgeable dietician to sit down with the patient and his family and go over the patient's likes and dislikes and tailor make a weekly menu with sufficient variety that will keep the patient satisfied. Many companies now have available a variety of low protein flour, recipes and ready made foods for these patients. When this is done, the diet can be quite tolerable despite its restrictions and most patients will abide by it because it makes them feel better. Most failures on the diet under these circumstances are due to progression of the renal disease to a point where diet alone cannot manage the symptoms and dialysis becomes necessary. This usually occurs at creatinine clearances less than 3 cc per minute.

So far the emphasis has been on protein restriction. But it is extremely important to remember that the patient must get sufficient protein in the diet to maintain nitrogen balance. Otherwise he will break down his own body protein and will get a clinical picture similar to Kwashiorkor^{8,9} with increased susceptibility to infection, cardiomyopathy and a rapid downhill course. Therefore patients who have excessive losses of protein such as nephrotics or patients undergoing dialysis, catabolic patients and patients on broad spectrum antibiotics may require greater amounts of protein in the diet to maintain nitrogen balance.^{1,10}

Carbohydrates If the Giovannetti diet is to maintain good nitrogen balance, it must provide adequate calories in the form of carbohydrates. The carbohydrates prevent the breakdown of body protein as an energy

source and, by stimulating insulin secretion, enhance the proper utilization of the essential amino acids. Two well known clinical observations that at first seem paradoxical but are readily explainable are, firstly, that diabetic patients who are on insulin or oral hypoglycemic agents require less of these medications after they develop renal failure, and secondly, that non diabetic patients who develop renal failure have a very high incidence of glucose intolerance resembling that seen in maturity onset diabetes. Since about 50 per cent of the plasma insulin is used and degraded by the kidney,¹¹ the insulin half life is prolonged in patients with renal failure thus explaining the decrease in insulin requirements in diabetic patients who develop renal failure. It has also been demonstrated that there is a peripheral insensitivity to endogenous insulin in patients with renal failure,^{12,13} thus explaining the abnormal glucose tolerance in non diabetic patients who develop renal failure.

These abnormalities in carbohydrate metabolism have some important clinical implications. The physician should be aware of the potential hazard of hypoglycemia in any diabetic patient who is on a fixed dose of hypoglycemic agents in whom renal insufficiency develops. In the emergency treatment of hyperkalemia the normal dose of insulin/glucose given intravenously is 1 unit of regular insulin per 4 Gm glucose. To avoid delayed hypoglycemic reactions secondary to the prolonged half life of insulin in patients with severe renal insufficiency, the insulin/glucose ratio should be 1 unit of insulin per 10 Gm glucose.¹⁴ Rarely rebound hypoglycemia and occasionally severe hyperglycemia with hyperosmolar coma have been reported in patients undergoing dialysis.^{15,16}

Lipids Hyperlipemia is a well known consequence of the nephrotic syndrome with or without renal failure. More recently it has been shown that patients with non nephrotic uremia also have hypertriglyceridemia.^{17,18} The exact mechanism for this is unknown but may be related to decreased tissue lipoprotein lipase and triglyceride removal capacity.¹⁹ Hemodialysis seems to accentuate the hyperlipemia both acutely and chronically.^{20,21} In general dietary

control and anti hyperlipidemic drugs are ineffective but a trial of both regimens should be attempted in every patient

Fluids Any patient with renal failure should receive the maximum amount of fluid per 24 hours that his kidneys can excrete without developing evidence of congestion or edema. The reason for this is that urea and possibly other toxic metabolic products back diffuse into the blood throughout the length of the tubules. An increase in urine flow through the tubule (increase in free water clearance) will increase the amount of these products excreted, since they would have less time to back diffuse into the blood.¹¹

To determine the ideal amount of fluid per 24 hours for any given patient body weight and fluid intake and output must be recorded daily. The physical exam and frequent monitoring of vital signs are extremely important in following the patient's state of hydration.¹ Changes in heart size on serial chest x rays are quite valuable. Where the signs and symptoms are equivocal a blood volume determination may be helpful. Once it is felt by all the clinical and other diagnostic parameters that the patient is at his ideal state of hydration the total fluids per 24 hours to be given equals the maximum urine output plus any other measurable losses plus 400 to 600 cc for insensible fluid loss.¹²

About 50 per cent of patients with renal failure on the Giovannetti diet retain water in spite of all restrictions placed on them so diuretics need to be used. Because of their very low glomerular filtration rates only the very potent diuretics such as furosemide (Lasix) will be effective. The dose can be increased until the desired diuretic effect is achieved which may be as much as 2 to 3 Gm of furosemide per day in divided doses.¹³

It is extremely important to stress the fact that there is no indication for digitalis in patients with congestion secondary to hypervolemia alone in renal failure.¹ If anything this drug is contraindicated in the patients because of its accumulation to toxic levels in renal failure and because of the marked fluctuations in serum potassium that can occur in the course of treating these patients with diuretics and dialy-

sis. Fluid restriction, diuretics and dialysis are the only proven safe methods of treating the congested state in these patients.¹ When digitalis must be used in patients with advanced renal failure (for therapy of severe arrhythmias or associated heart failure that cannot be managed by control of fluid volume alone) digitoxin is the drug of choice.²⁴

Sodium Unlike patients with congestive heart failure who always tend to have severe sodium retention excreting less than 10 mEq per liter of urine patients with renal failure tend to lose more sodium. This will vary from patient to patient at the same glomerular filtration rate depending on the underlying renal disease—i.e. patients with tubular involvement such as pyelonephritis will lose more sodium than patients with glomerulonephritis. The sodium loss will also vary in the same patient at different levels of renal function. Therefore since the patient's kidneys have lost their ability to significantly regulate the amount of sodium excreted per 24 hours they are all potentially salt losers or salt retainers depending on whether the dietary sodium is less or greater than the amount excreted. The implications of these are obvious: too much sodium in the diet can contribute to or aggravate the patient's hypertension and congestion; too little sodium may result in muscle cramps, convulsions, hypovolemia, hypotension and further deterioration of renal function.¹¹ To calculate the amount of sodium to be allowed in the diet one needs to know the 24 hour urinary sodium loss. If renal function is not stable as in the diuretic phase of acute renal failure or if there are extrarenal losses of sodium such as vomiting the 24 hour urinary sodium and other sodium losses must be measured every day and the intake adjusted accordingly.

The treatment of sodium deficiency is easy with increase of replacement therapy. In patients who are sodium retainers diuretics may have to be used. Salt substitutes need careful handling in renal failure patients since the cation substituted is potassium and each packet (1 Gm weight) has about 12 mEq potassium.¹²

Potassium Hyperkalemia is a common problem in severe renal insufficiency often aggravated by the associated acidosis. The

amount of potassium to be allowed in the diet in any given patient can be estimated by the 24 hour urinary potassium. If, despite all restriction the patient still tends to be hyperkalemic, exchange resins can be given orally. If a sodium exchange resin (exchanging about 2 to 3 mEq sodium per Gm resin²⁵) is used, one may run into problems with sodium retention and edema. Calcium and aluminum resins have also been used but the former can cause hypercalcemia^{26, 27} and aluminum intoxication has been reported with the latter.^{27, 28}

The emergency treatment of life threatening hyperkalemia^{25, 26} consists of intravenous glucose and insulin (10 Gm glucose per 1 unit regular insulin), calcium sodium bicarbonate and hypertonic saline solution. Since most patients with renal failure are hypovolemic sodium bicarbonate and saline cannot be given safely. In general I have found that 5 units of regular insulin given intravenously followed by a bolus of 50 c.c. of 50 per cent dextrose and an intravenous drip of 500 c.c. of 10 per cent dextrose in water containing 5 Gm of calcium gluconate run at 100 c.c. per hour will bring the serum potassium down immediately and protect the patient from the cardiovascular effects of hyperkalemia until the oral exchange resins or enemas begin to take effect.¹²

Hypokalemia may be seen at any stage of renal failure and should be treated because of its long term adverse effects on the kidney. But the replacement therapy must be slow and serum potassium must be carefully monitored since it takes several hours for the extracellular potassium to come into complete equilibration with the intracellular pool.¹² If potassium sparing diuretics are to be used potassium supplements must be stopped.¹

Acidosis Most patients with severe renal failure have some degree of compensated metabolic acidosis. The use of the oral phosphate binding gels tends to partially alleviate this. Since sodium and magnesium may be retained in renal failure antacids that are low in these cations are preferred. Some workers have used calcium carbonate²⁹ as an antacid in those patients who have a low serum calcium and severe renal osteodystrophy. The main complication of this form of therapy is hypercalcemia.³¹

In severe decompensated metabolic acidosis, intravenous sodium bicarbonate may be used if the patient can tolerate the sodium load. In patients with extensive tubular involvement that wastes a great deal of bicarbonate sodium in the diet can be restricted and sodium bicarbonate tablets can be used as supplements for control of the acidosis.¹²

Since lactate is normally converted to bicarbonate by the liver one must be careful in the use of any intravenous or peritoneal dialysis solutions which contain lactate as the main buffer in patients with associated lactic acidosis and/or severe liver disease. It is contraindicated in the former and when it is used in patients with liver disease arterial pH should be checked early in the course of therapy to ensure the acidosis is being corrected rather than made worse.¹²

Minerals and vitamins Since the dietary intake may vary in patients with renal failure, and since water soluble vitamins may be removed during dialysis,^{1, 2, 32} the water soluble vitamins should be given as supplements to all patients with renal failure. Patients who are hypocalcemic with raised alkaline phosphatase and radiographic evidence of severe renal osteodystrophy may be tried on vitamin D or its more active metabolites such as 25 hydroxycholecalciferol.¹² But this is to be done under close supervision since hypercalcemia and severe soft tissue calcification may occur as a result of therapy.³³

Since the serum magnesium levels in most patients with advanced renal failure is normal or high irrespective of the intracellular magnesium one usually tries to avoid large doses of magnesium in the form of antacids or laxatives.¹²

In general the anemia of patients with renal failure is not associated with significant iron deficiency in spite of the poor absorption of oral iron in these patients.¹² But some patients on chronic hemodialysis become iron deficient because of frequent venepuncture and blood loss in the coil and here supplemental iron therapy becomes necessary.³⁴

I would like to thank Dr. Albert Rubin and Dr. Paul Bienstock for their advice and guidance in preparing the manuscript.

REFERENCES

- 1 Domenech J G and Evans D W Uremia in congestive heart failure *Quart J Med* 28 117 1969
- 2 Brown C L Hill M J and Richards P Bacterial ureasemia in uremic man *Lancet* 2 406 1971
- 3 Giordano C DePascale C Balestrieri C Cattadini D and Crescenzi A Incorporation of urea N¹⁵ in amino acids of patients with chronic renal failure on low nitrogen diet *Am J Clin Nutr* 21 394 1968
- 4 Berlyne G M and Hocken A G The dietary treatment of chronic renal failure in Berlyne G M ed Nutrition in renal disease Baltimore 1968 The Williams & Wilkins Company pp 33-54
- 5 Hersted D M Minimum protein requirements of adults *Am J Clin Nutr* 21 352 1968
- 6 Blainey J D and Chamberlain M J Dietary treatment of chronic renal failure *Br Med Bull* 27 160 1971
- 7 Berlyne G M Gaan D and Ginks W R Dietary treatment of chronic renal failure *Am J Clin Nutr* 21 547 1968
- 8 Metcalf J Ionic composition and cell metabolism with protein-calorie malnutrition in man *Am J Clin Nutr* 21 376 1968
- 9 Holt I E Jr Some problems in dietary amino acids requirements *Am J Clin Nutr* 21 367 1968
- 10 Bailey G L Hampers C L and Merrill J P Reversible cardiomyopathy in uremia *Trans Am Soc Artif Intern Organs* 13 263 1967
- 11 Wharton B A Balmer S E Somers K and Templeton A C The myocardium in kwashiorkor *Quart J Med* 38 107 1969
- 12 David D S Hochgelerant E Rubin A I and Stenzel K H Dietary management in renal failure *Lancet* 2:34 1972
- 13 Ginn H E Frost A and Lacey W W Nitrogen balance in hemodialysis patients *Am J Clin Nutr* 21 385 1968
- 14 O'Brien J P and Sharpe A R Jr The influence of renal disease on the insulin¹²⁵I appearance in man *Metabolism* 16:16 1967
- 15 Hampers C L Soeldner J S Gleason R E Bailey G L Diamond J A and Merrill J P Insulin-glucose relationships in uremia *Am J Clin Nutr* 21:414 1968
- 16 Spatz I M Rubenstein A H Berison I Abrahams C and Lowy C Carbohydrate metabolism in renal disease *Quart J Med* 39:101 1970
- 17 Rige G A and Bercu B A Hypoglycemia—a complication of hemodialysis *N Engl J Med* 277:1139 1967
- 18 Porter D J Death as a result of hyperglycemia without ketoacidosis: a complication of hemodialysis *Ann Intern Med* 65:399 1966
- 19 Boyer J Gill G N and Epstein F H Hypoglycemia and hyperosmolality complicating peritoneal dialysis *Ann Intern Med* 67:568 1967
- 20 Bagdade J D Ponte D Jr and Bierman E L Hypertriglyceridemia—a metabolic consequence of chronic renal failure *N Engl J Med* 279:181 1968
- 21 Bagdade J D Lipemia a sequel of chronic renal failure and hemodialysis *Am J Clin Nutr* 21 426 1968
- 22 Saltas T T and Friedman E A Plasma lipids studies of uremic patients during hemodialysis *Am J Clin Nutr* 21 430 1968
- 23 Fleisher D S Vocci G Garfunkel J Peruganan H Kirkpatrick J Jr Wells C R and McElfresh A F Hemodynamics findings in acute glomerulonephritis *J Pediatr* 69 1054 1966
- 24 Rasmussen K Jervell J Storstein L and Gjerdrum K Digoxin kinetics in patients with impaired renal function *Clin Pharmacol Ther* 13 6 1971
- 25 Papadimitriou M Gingell J C and Chisholm G D Hypercalcemia from calcium ion exchange resin in patients on regular hemodialysis *Lancet* 2 948 1968
- 26 Sevitt L H and Wrong O M Hypercalcemia from calcium resin in patients with chronic renal failure *Lancet* 2 950 1968
- 27 Berlyne G M Yagil R Ari J B Weinberger G Knopf E and Danovitch G M Aluminum toxicity in rats *Lancet* 1 564 1972
- 28 Thurston H Gilmore G R and Swales J D Aluminum retention and toxicity in chronic renal failure *Lancet* 1:881 1972
- 29 Berlyne G M and Yagil R Aluminum toxicity *Lancet* 2:47 1972
- 30 Makoff D L Gordon A Franklin S S Gerstein A R and Maxwell M H Chronic calcium carbonate therapy in uremia *Arch Intern Med* 123 15 1969
- 31 Stiel J N Mitchell C A Radcliff F J and Piper D W Hypercalcemia in patients with peptic ulceration receiving large doses of calcium carbonate *Gastroenterology* 53 900 1967
- 32 Sullivan J F and Eisenstein A B Ascorbic acid depletion in patients undergoing chronic hemodialysis *Am J Clin Nutr* 23 1339 1970
- 33 Lasker N Harvey A and Baker H Vitamin levels in hemodialysis and intermittent peritoneal dialysis *Trans Am Soc Artif Intern Organs* 9 51 1963
- 34 Clinco-pathologic conference *Am J Med* 41:593 1966
- 35 Hampers C I Stessl R Nathan D Snyder D and Merrill J P Megaloblastic hematopoiesis in uremia and in patients on long term hemodialysis *N Engl J Med* 276:551 1967
- 36 Mallick N P and Berlyne G M Arterial calcification after vitamin D therapy in hyperphosphatemic renal failure *Lancet* 2 1316 1968

amount of potassium to be allowed in the diet in any given patient can be estimated by the 24 hour urinary potassium. If, despite all restriction, the patient still tends to be hyperkalemic exchange resins can be given orally. If a sodium exchange resin (exchanging about 2 to 3 mEq sodium per Gm resin²²) is used, one may run into problems with sodium retention and edema. Calcium and aluminum resins have also been used, but the former can cause hypercalcemia^{23, 24} and aluminum intoxication has been reported with the latter²⁵.

The emergency treatment of life threatening hyperkalemia^{26, 27} consists of intravenous glucose and insulin (10 Gm glucose per 1 unit regular insulin), calcium sodium bicarbonate and hypertonic saline solution. Since most patients with renal failure are hypovolemic sodium bicarbonate and saline cannot be given safely. In general I have found that 5 units of regular insulin given intravenously followed by a bolus of 50 c.c. of 50 per cent dextrose and an intravenous drip of 500 c.c. of 10 per cent dextrose in water containing 5 Gm of calcium gluconate run at 100 c.c. per hour will bring the serum potassium down immediately and protect the patient from the cardiovascular effects of hyperkalemia until the oral exchange resins or enemas begin to take effect¹².

Hypokalemia may be seen at any stage of renal failure and should be treated because of its long term adverse effects on the kidney. But the replacement therapy must be slow and serum potassium must be carefully monitored since it takes several hours for the extracellular potassium to come into complete equilibration with the intracellular pool²⁸. If potassium sparing diuretics are to be used potassium supplements must be stopped¹².

Acidosis Most patients with severe renal failure have some degree of compensated metabolic acidosis. The use of the oral phosphate binding gels tends to partially alleviate this. Since sodium and magnesium may be retained in renal failure antacids that are low in these cations are preferred. Some workers have used calcium carbonate²⁹ as an antacid in those patients who have a low serum calcium and severe renal osteodystrophy. The main complication of this form of therapy is hypercalcemia³¹.

In severe decompensated metabolic acidosis, intravenous sodium bicarbonate may be used if the patient can tolerate the sodium load. In patients with extensive tubular involvement that wastes a great deal of bicarbonate, sodium in the diet can be restricted and sodium bicarbonate tablets can be used as supplements for control of the acidosis³².

Since lactate is normally converted to bicarbonate by the liver one must be careful in the use of any intravenous or peritoneal dialysis solutions which contain lactate as the main buffer in patients with associated lactic acidosis and/or severe liver disease. It is contraindicated in the former and when it is used in patients with liver disease arterial pH should be checked early in the course of therapy to ensure the acidosis is being corrected rather than made worse³³.

Minerals and vitamins Since the dietary intake may vary in patients with renal failure, and since water soluble vitamins may be removed during dialysis^{1, 32, 33} the water soluble vitamins should be given as supplements to all patients with renal failure. Patients who are hypocalcemic with raised alkaline phosphatases and radiographic evidence of severe renal osteodystrophy may be tried on vitamin D or its more active metabolites such as 25 hydroxycholecalciferol³⁴. But this is to be done under close supervision since hypercalcemia and severe soft tissue calcification may occur as a result of therapy³⁵.

Since the serum magnesium levels in most patients with advanced renal failure is normal or high irrespective of the intracellular magnesium, one usually tries to avoid large doses of magnesium in the form of antacids or laxatives³⁶.

In general the anemia of patients with renal failure is not associated with significant iron deficiency in spite of the poor absorption of oral iron in these patients¹. But some patients on chronic hemodialysis become iron deficient because of frequent venepuncture and blood loss in the coil and here supplemental iron therapy becomes necessary³⁷.

I would like to thank Dr. Albert Rubin and Dr. Paul Biensack for their advice and guidance in preparing the manuscript.

participant in the study. In addition all infants with congenital heart disease who died and who did not have cardiac catheterization or cardiac surgery also entered the study. A total of 262 infants satisfied these criteria in the years 1969 and 1970.

Each of the 5 diagnostic centers was visited by one of the authors (George Kelso). Patients were identified from catheterization autopsy or surgical log books and from card indexes in the medical record libraries. More detailed information was then obtained from cardiac catheterization surgical and autopsy reports. Follow up information was abstracted from hospital records. The total births for the years 1969 and 1970 was furnished by the State Department of Health. All patients were also identified by county of residence. The data were punched on 50 column IBM computer cards and were processed by means of a card sorter.

Results

A Incidence The total number of infants entering the study was 262 giving a calculated incidence of 1.7/1 000 births. The predicted incidence at 2.1/1 000 births is 304 infants. Therefore 86 per cent of the infants estimated to be at risk were identified in the 5 centers. Forty two per cent of the infants were cared for at Milwaukee Children's Hospital, 29 per cent at St Mary's Hospital, 18 per cent at University Hospital, 6 per cent at Gunderson Clinic and 5 per cent at Marshfield Clinic. Eighty nine per cent of the infants were seen at hospitals closely affiliated with a medical school. This information is shown in Table I. Table I also gives the criteria used to admit a patient to the study (catheterization surgery without prior catheterization or autopsy). It is seen that 73 per cent (192) of the infants were identified primarily by cardiac catheterization. Twenty one per cent (53) of the patients died prior to any diagnostic or therapeutic intervention and were admitted to the study on the basis of autopsy findings. Seventeen patients had surgery without any prior catheterization. All of these infants were from the same institution and 16/17 were premature infants who had ligation of a patent ductus arteriosus.

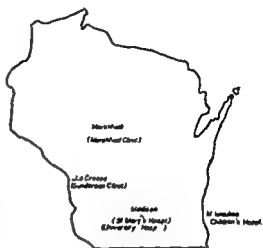


Fig 1 Map of Wisconsin giving the locations of the 5 centers caring for infants with critical congenital heart disease.

B Frequency of specific lesions The most commonly encountered anomaly was ventricular septal defect (VSD) (19 per cent) followed by transposition of the great arteries (TGA) (13 per cent) and patent ductus arteriosus (PDA) (11 per cent). Other frequently seen lesions were hypoplastic left heart syndrome, tetralogy of Fallot, coarctation of the aorta, total anomalous pulmonary venous drainage and pulmonary valve obstructions—stenosis and atresia. More than one fifth of the patients fell into the other category which included the less common problems: patients with multiple congenital anomalies and genetic disorders. In those patients where more than one lesion existed the most important anatomic abnormality was used as a basis for the classification. This material is summarized in Table II.

C Age first identified The age range at which these infants were first identified is given in Table III. Twenty seven per cent (69) were seen during the first week of life, forty eight per cent (125) by the first month and seventy per cent (182) by the first 3 months. By 3 months of age 87 per cent of the patients had been identified. Certain lesions resulted in significant signs and symptoms earlier than others. Thus 93 per cent of the patients with hypoplastic left heart syndrome, 61 per cent of those with transposition, 58 per cent of the infants with aortic coarctation and 54 per

Demography of critical congenital heart disease

George F. Kelso, B.S.*

William J. Gallen, M.D.

David Z. Friedberg, M.D.

Milwaukee, Wisc

Aggressive medical and surgical intervention is the accepted mode of therapy for critically ill infants with congenital heart disease.^{1,2} Without such intervention it has been demonstrated that 2/1 000 live born infants will die during the first year of life due to congenital cardiac anomalies.^{4,5} This group of infants can be considered to have "critical congenital heart disease." The salvage of these infants thus becomes dependent on early identification (usually in the neonatal period) and subsequent referral to centers having specialized facilities for diagnosis and treatment.

The State of Wisconsin, with a population of 4,417 731 (1970 census) has a large urban area in its southeast portion (Milwaukee and vicinity) but is otherwise composed of rural farming areas. Only five medical facilities in this state care for infants with congenital heart disease. They are (1) Milwaukee Children's Hospital, Milwaukee—the pediatric teaching unit of The Medical College of Wisconsin; (2) University Hospital, Madison—the pediatric teaching unit of The University of Wisconsin Medical School; (3) St. Mary's Medical Center, Madison—a private insti-

tution affiliated with The University of Wisconsin; (4) Marshfield Clinic, Marshfield—a large multi-specialty private clinic; and (5) Gunderson Clinic, La Crosse—in other large multi-specialty private clinic. The location of these centers is seen on the map in Fig. 1. As far as can be determined, no other facility in the state cares for infants with critical congenital heart disease.

It was the purpose of this study (1) to identify all infants with critical congenital heart disease who were born in Wisconsin in 1969 and 1970 and to compare this incidence with that predicted from the literature; (2) to provide information on the frequency of specific lesions on the age of the patient at the time of diagnostic or therapeutic intervention; the mode of therapy (medical or surgical) and the final outcome of this group of infants; (3) to follow these children for at least 1 year; and (4) to analyze the referral patterns of these infants in the State of Wisconsin.

Material and methods

Any infant under 1 year of age who had congenital heart disease and whose clinical state was such that cardiac catheterization and/or surgery was performed became a

From the Fenchel Cardiac Study Center, Milwaukee Children's Hospital and The Department of Pediatrics, The Medical College of Wisconsin, Milwaukee, Wisc.

Received for publication Aug. 16, 1972.

Reprint requests to David Z. Friedberg, M.D., Fenchel Cardiac Study Center, Milwaukee Children's Hospital,

1700 W. Wisconsin Ave., Milwaukee, Wisc. 53233.

*This study was done while Mr. Kelso was a fourth year student at The Medical College of Wisconsin.

participant in the study. In addition all infants with congenital heart disease who died and who did not have cardiac catheterization or cardiac surgery also entered the study. A total of 262 infants satisfied these criteria in the years 1969 and 1970.

Each of the 5 diagnostic centers was visited by one of the authors (George Helko). Patients were identified from catheterization autopsy or surgical log books and from card indexes in the medical record libraries. More detailed information was then obtained from cardiac catheterization surgical and autopsy reports. Follow up information was abstracted from hospital records. The total births for the years 1969 and 1970 was furnished by the State Department of Health. All patients were also identified by county of residence. The data were punched on 50 column IBM computer cards and were processed by means of a card sorter.

Results

A Incidence The total number of infants entering the study was 262 giving a calculated incidence of 1.7/1 000 births. The predicted incidence at 2/1 000 births is 304 infants. Therefore 86 per cent of the infants estimated to be at risk were identified in the 5 centers. Forty two per cent of the infants were cared for at Milwaukee Children's Hospital, 29 per cent at St. Mary's Hospital, 18 per cent at University Hospital, 6 per cent at Gunderson Clinic and 5 per cent at Marshfield Clinic. Eighty nine per cent of the infants were seen at hospitals closely affiliated with a medical school. This information is shown in Table I. Table I also gives the criteria used to admit a patient to the study (catheterization surgery without prior catheterization or autopsy). It is seen that 73 per cent (197) of the infants were identified primarily by cardiac catheterization. Twenty one per cent (53) of the patients died prior to any diagnostic or therapeutic intervention and were admitted to the study on the basis of autopsy findings. Seventeen patients had surgery without any prior catheterization. All of these infants were from the same institution and 16/17 were premature infants who had ligation of a patent ~~ductus~~ ^{aorta} ~~arteriosus~~.



Fig. 1 Map of Wisconsin giving the locations of the 5 centers caring for infants with critical congenital heart disease.

B Frequency of specific lesions The most commonly encountered anomaly was ventricular septal defect (VSD) (19 per cent) followed by transposition of the great arteries (TGA) (13 per cent) and patent ductus arteriosus (PDA) (11 per cent). Other frequently seen lesions were hypoplastic left heart syndrome, tetralogy of Fallot, coarctation of the aorta, total anomalous pulmonary venous drainage and pulmonary valve obstructions—stenosis and atresia. More than one fifth of the patients fell into the 'other' category which included the less common problems: patients with multiple congenital anomalies and genetic disorders. In those patients where more than one lesion existed the most important anatomic abnormality was used as a basis for the classification. This material is summarized in Table II.

C Age first identified The age range at which these infants were first identified is given in Table III. Twenty seven per cent (69) were seen during the first week of life, forty-eight per cent (125) by the first month and seventy per cent (182) by the first 3 months. By 6 months of age 87 per cent of the patients had been identified. Certain lesions resulted in significant signs and symptoms earlier than others. Thus 93 per cent of the patients with hypoplastic left heart syndrome, 61 per cent of those with transposition, 58 per cent of the infants with aortic coarctation and 54 per

Demography of critical congenital heart disease

George F Kelso, B S*

William J Gallen, M D

David Z Friedberg, M D

Milwaukee, Wisc

Aggressive medical and surgical intervention is the accepted mode of therapy for critically ill infants with congenital heart disease.^{1,2} Without such intervention, it has been demonstrated that 2/1 000 live born infants will die during the first year of life due to congenital cardiac anomalies.^{3,4} This group of infants can be considered to have "critical congenital heart disease." The salvage of these infants thus becomes dependent on early identification (usually in the neonatal period) and subsequent referral to centers having specialized facilities for diagnosis and treatment.

The State of Wisconsin with a population of 4,417,731 (1970 census) has a large urban area in its southeast portion (Milwaukee and vicinity) but is otherwise composed of rural farming areas. Only five medical facilities in this state care for infants with congenital heart disease. They are (1) Milwaukee Children's Hospital Milwaukee—the pediatric teaching unit of The Medical College of Wisconsin (2) University Hospital Madison—the pediatric teaching unit of The University of Wisconsin Medical School (3) St Mary's Medical Center, Madison—a private insti-

tution affiliated with The University of Wisconsin, (4) Marshfield Clinic, Marshfield—a large multi specialty private clinic, and (5) Gunderson Clinic La Crosse—an other large multi specialty private clinic. The location of these centers is seen on the map in Fig 1. As far as can be determined no other facility in the state cares for infants with critical congenital heart disease.

It was the purpose of this study (1) to identify all infants with critical congenital heart disease who were born in Wisconsin in 1969 and 1970 and to compare this incidence with that predicted from the literature, (2) to provide information on the frequency of specific lesions on the age of the patient at the time of diagnostic or therapeutic intervention, the mode of therapy (medical or surgical) and the final outcome of this group of infants (3) to follow these children for at least 1 year and (4) to analyze the referral patterns of these infants in the State of Wisconsin.

Material and methods

Any infant under 1 year of age who had congenital heart disease and whose clinical state was such that cardiac catheterization and/or surgery was performed became a

From the Fairchild Cardiac Study Center Milwaukee Children's Hospital and The Department of Pediatric The Medical College of Wisconsin Milwaukee Wisc

Received for publication Aug 16 1972

Reprint requests to David Z Friedberg M D Fairchild Cardiac Study Center Milwaukee Children's Hospital 1700 W Wisconsin Ave Milwaukee Wisc 53233

*This study was done while Mr Kelso was a fourth year student at The Medical College of Wisconsin

Table III Age at which lesion was first identified in 262 patients

| LESION | NUMBER OF PATIENTS | <1 WK | 1-4 WKS | 1-3 MOS | 3-6 MOS | 6-12 MOS |
|-----------------------------|--------------------|---------|---------|---------|---------|----------|
| VSD | 49 | 5 | 11 | 12 | 14 | 7 |
| Transposition | 33 | 11 | 9 | 8 | 1 | 4 |
| Potential Ductus Arteriosus | 28 | 5 | 9 | 6 | 5 | 2 |
| Hypoplastic Left Heart | 26 | 20 | 5 | 0 | 1 | 0 |
| Tetralogy of Fallot | 23 | 2 | 5 | 6 | 6 | 4 |
| Coarctation of Aorta | 19 | 4 | 7 | 1 | 4 | 3 |
| Total Anomalous Pul Vn | 11 | 1 | 1 | 6 | 2 | 1 |
| Pulmonary Atresia | 7 | 3 | 0 | 3 | 0 | 1 |
| Pulmonary Stenosis | 7 | 2 | 2 | 1 | 0 | 2 |
| Other | 59 | 16 | 7 | 14 | 12 | 10 |
| TOTAL | 262 | 69(27%) | 56(21%) | 57(22%) | 46(17%) | 34(13%) |

Table IV Number and per cent of patients surviving out of 262 patients

| LESION | No PTS | No PTS T AUT | No SURVIVING | % SURVIVING | % SURVIVING T AUT |
|--------------------|--------|--------------|-------------------------------------|-------------|-------------------|
| Vent Septal Defect | 49 | 42 | 40 ^{M 17} _{S 23} | 82 | 95 |
| Transposition | 33 | 29 | 12 ^{M 6} _{S 6} | 36 | 41 |
| Potential Ductus | 28 | 25 | 21 ^{M 1} _{S 20} | 75 | 84 |
| Hypoplastic LH | 26 | 11 | 0 ^{M 0} _{S 0} | 0 | 0 |
| Tetralogy | 23 | 21 | 14 ^{M 8} _{S 6} | 61 | 67 |
| Coarctation | 19 | 16 | 12 ^{M 6} _{S 6} | 63 | 75 |
| Total An Pul Vn | 11 | 8 | 5 ^{M 3} _{S 2} | 45 | 53 |
| Pul Atresia | 7 | 6 | 2 ^{M 0} _{S 2} | 29 | 33 |
| Pul Stenosis | 7 | 7 | 7 ^{M 2} _{S 5} | 100 | 100 |
| Other | 59 | 44 | 26 ^{M 19} _{S 7} | 45 | 60 |
| TOTAL | 262 | 209 | 139 ^{M 64} _{S 75} | 62 54% | 62 66% |

M Medical Follow

S Surgical Follow

died. Fifty three of these died before cardiac catheterization or surgery was performed. Thirteen patients died within 24 hours of cardiac catheterization. This represented a total catheterization mortality rate of 63 per cent. Approximately equal numbers of patients who were fol-

lowed medically (25) and surgically (32) succumbed. Results are summarized in Table V.

F Surgery There were 109 surgical procedures. In those situations where more than one surgical procedure was done during the same operation, the primary pro-

Table 1 Incidence of critical congenital heart disease

| INSTITUTION | No PATIENTS | CRITERIA FOR IDENTIFICATION | | |
|--------------------|-------------|-----------------------------|----------|---------|
| | | CATH | AUTOPSY | SURGERY |
| Milw Child Hosp | 112 (42%) | 89 | 33 | — |
| St Mary's Hosp | 77 (29%) | 49 | 11 | 17 |
| University Hosp | 44 (18%) | 33 | 11 | — |
| Gunderson Clinic | 18 (6%) | 11 | 4 | — |
| Marshfield Clinic | 14 (5%) | 10 | 4 | — |
| TOTAL | 262 | 192 (73%) | 53 (21%) | 17 (6%) |
| PREDICTED (2/1000) | 304 | | | |

Table 11 Frequency of specific lesions in 262 patients

| LESION | NUMBER OF PATIENTS | PERCENT OF TOTAL |
|---------------------------------|--------------------|------------------|
| Ventricular Septal Defect | 49 | 19% |
| Transposition | 33 | 13% |
| Patent Ductus Arteriosus | 28 | 11% |
| Hypoplastic Left Heart | 26 | 10% |
| Tetralogy of Fallot | 23 | 9% |
| Coarctation of Aorta | 19 | 7% |
| Total Anomalous Pul Vn | 11 | 4% |
| Pulmonary Atresia-Intact Septum | 7 | — |
| Pulmonary Stenosis | 7 | — |
| Other | 59 | 22% |
| TOTAL | 262 | |

cent having patent ductus arteriosus were seen during the first month of life. This was in contrast to patients with tetralogy of Fallot, total anomalous pulmonary venous return, and ventricular septal defect—the majority of whom were identified after the first month of life.

D. Survival At the end of at least 1 year of follow up 54 per cent of patients (139) were alive. However, if we eliminate the 53 patients who entered the study by autopsy without the benefit of catheterization and further surgical therapy, the survival rate rises to 66 per cent. Of the 139 patients surviving 45 per cent (62) were treated medically while 55 per cent

(77) had surgery. The best salvage was seen in patients with pulmonary stenosis (100 per cent) VSD (82 per cent) PDA (75 per cent) coarctation (63 per cent) and tetralogy of Fallot (61 per cent). If patients entering the study by autopsy are eliminated, then salvage rates go up to 95 per cent in VSD, 84 per cent in PDA, 75 per cent in coarctation and 67 per cent in tetralogy of Fallot. No patients with hypoplastic left heart survived—none had surgery. The salvage in transposition of the great arteries was disappointing (36 per cent). This information is seen in Table IV.

E. Deaths At the end of 1 year of follow up 46 per cent of the patients (123) had

referral thus becomes obvious. It is in this area that we feel progress can be made in reducing the over all mortality rate of infants with critical congenital heart disease.

The frequency of specific lesions agrees quite well with that of other studies.^{2,6,7} The small variations are in part due to the fact that we chose to examine the first year of life while others^{2,7} have concentrated on the first month only. This would obviously skew the diagnostic incidence towards the lesions whose life threatening characteristics became apparent earlier—the transposition pulmonary atresia and aortic coarctation. As has been mentioned the increased number of patients with patent ductus arteriosus relates to the fact that one institution was very interested in the ductus arteriosus of the premature and tended to be more aggressive in its diagnosis and treatment. If these patients were excluded patent ductus arteriosus would then comprise only 6 per cent of the total.

In a recent study by Feldt and associates¹ 43.5 per cent of the patients were diagnosed within the first week of life as opposed to 27 per cent of our patients. The Feldt study only included the surrounding county whereas we were concerned with an entire state. It is quite possible that the distance from hospital of birth to referring center was sufficient to delay referral. Again this time lag from place of birth to referral center may explain the large number of patients in this study who succumbed prior to any diagnostic or therapeutic intervention.

The total salvage rate was 54 per cent. This compares favorably with other studies: Coleman⁴ 42 per cent, Varghese and colleagues² 43 per cent, Fyler and associates⁶ 64 per cent. If we exclude all infants who died prior to catheterization or surgery the salvage rate is increased to 69 per cent. Essentially then earlier referral would result in a much higher survival rate in these patients. The surgical survival rate was 71 per cent while 41 per cent of those patients treated medically were alive at the end of 1 year. This confirms the opinion of Fyler and colleagues⁶ who feel that aggressive management including surgery produces the best results. Again the over all surgical results are similar to those of others.¹

Approximately 40 per cent of the surgical procedures were palliative in nature and further surgical intervention in these infants is anticipated in the future. A certain number of salvaged infants will probably not survive the second operation. It is expected that recent trends towards early total correction using deep hypothermia with cardiac arrest¹⁰ will decrease the necessity for repeated high risk surgery.

It is thus seen that early referral accurate diagnosis and aggressive surgical intervention can contribute to salvage of a significantly large group of infants who were only recently thought to be incurable.¹¹ The burden thus falls on the primary physician to recognize the presence of critical congenital heart disease and institute appropriate referral and on the cardiac center to have available on a 24 hour per day basis the facilities for anatomical diagnosis and treatment. The need for rapid and safe transportation systems and excellent communication is obvious. We feel strongly that no hospital should assume responsibility for care of these infants unless all of the necessary facilities and personnel are available and that the care of these infants requires a complete and total commitment on the part of those assuming this responsibility.

Summary

In a retrospective study the incidence of critical congenital heart disease in the State of Wisconsin was determined for the years 1969-1970. Two hundred sixty two infants were identified in 5 pediatric cardiology centers giving an incidence of 1.7 cases per 1000 live births. Patients were admitted to the study on the basis of cardiac catheterization (192), autopsy (53) or surgery without prior catheterization (17).

Lesions most commonly encountered were ventricular septal defects (19 per cent), transposition of the great arteries (13 per cent), patent ductus arteriosus (11 per cent), hypoplastic left heart (10 per cent) and tetralogy of Fallot (9 per cent). Analysis was made to relate the specific cardiac anomaly to the age at which diagnostic and/or therapeutic intervention was required.

One hundred nine surgical procedures were performed. These included pul-

Table V Number and per cent of deaths in 262 patients

| PATIENTS DEAD = 123 (46 %) | | |
|----------------------------|----|--------|
| DEAD BEFORE CATH— | 53 | (44 %) |
| DEAD DUE TO CATH— | 13 | (10 %) |
| DEAD — SURGICAL — | 32 | (26 %) |
| DEAD — MEDICAL — | 25 | (20 %) |

Table VI Surgical results

| <u>PROCEDURE</u> | <u>NO PTS</u> | <u>ALIVE</u> | <u>DEAD</u> | <u>SURVIVAL %</u> |
|--------------------------|---------------|--------------|-------------|-------------------|
| Pulmonary Artery Band | 31 | 24 | 7 | 77 % |
| PDA Ligation | 24 | 20 | 4 | 83 % |
| Waterston Shunt | 18 | 8 | 10 | 44 % |
| Resection of Coarctation | 8 | 6 | 2 | 75 % |
| Atrial Septectomy | 8 | 5 | 3 | 63 % |
| Brock Procedure | 3 | 2 | 1 | 67 % |
| Other | 17 | 12 | 5 | 70 % |
| TOTAL | 109 | 77 | 32 | 71 % |

cedure was chosen for classification. Seventy one per cent of the patients (77) survived the surgery. The most successful procedures included pulmonary artery banding (77 per cent survived), ligation of a patent ductus arteriosus (83 per cent survived) and resection of coarctation of aorta (75 per cent survived). See Table VI.

G Referral patterns. All patients were identified as to county of residence. The center receiving a majority of patients from a particular county received credit as that county's referral area. Proximity determined referral area. In this sense, each of the 5 centers had its own naturally chosen referral population. Only in counties equal in distance between 2 or more centers was there a significant division in referral patterns. In general, it was both the parents and physicians desire that the infant be cared for closest to home.

Discussion

The incidence of critical congenital heart disease comprising 17/1 000 live births is somewhat lower than expected (2/1 000). However referrals to neighboring states have not been included and certainly some patients are referred out of state. The numbers leaving the state are probably not very significant as the referral patterns previously described, emphasize proximity as one of the most important factors in determining the center chosen for referral. Some of the undetected infants are thus dying without referral—probably undiagnosed as having congenital heart disease.

Of more concern is the fact that 21 per cent of all patients (53) died before catheterization or surgery could be accomplished and this represented 44 per cent of all deaths—a significant contribution to the total mortality rate. The net earlier

Complete right bundle branch block with left axis deviation: Significance of vectorcardiographic morphology

Edgar Lichstein MD
Kul D Chadda MD
Prem K Gupta MD
Elmhurst NY

Complete right bundle branch block with left axis deviation in the frontal plane was noted in 1934 by Wilson Johnston and Barker.¹ It was thought that this unusual form of block might in some cases be due to a lesion of one of the subdivisions of the left bundle in addition to right bundle branch block. Lenegre² and Unger and associates³ provided pathological support for this interpretation. Further confirmation was provided by the experimental animal studies of Watt and co-workers⁴ in which sectioning the right bundle branch after ligating the anterior fibers of the left bundle branch produced the electrocardiographic pattern in question.

Studies of vectorcardiographic morphology have attempted to find a relationship between pattern and clinical course. Salzman and colleagues⁵ described two basic vectorcardiographic loops which were characterized by the rotation in the horizontal plane. Patients with a Type A or counterclockwise horizontal loop had a better prognosis than patients with a Type B clockwise horizontal loop. A similar study

by Kulbertus and co-workers⁶ denied any significant difference between the two groups.

This study is concerned with a large group of patients with complete right bundle branch block and abnormal left axis deviation. An attempt is made to correlate the vectorcardiographic morphology and the His bundle electrogram with certain historical features such as dizziness and syncope.

Patients and method

The patients in this study included both inpatients and outpatients of a large municipal hospital. The majority of patients were discovered as a result of screening routinely requested ECGs. Other patients were seen in consultation either because of their underlying cardiac disease or for specific evaluation of an abnormal ECG. History and physical examination was performed by the authors at the time the vectorcardiogram (VCG) was recorded. Hypertension was thought to be present if repeated blood pressure determinations

From the Department of Medicine, Division of Cardiology, Mount Sinai Hospital Service, City Hospital Center, and Elmhurst Medical School, I Medical Center of the City University of New York.
Received for publication August 28, 1972.
Reprint requests to Edgar Lichstein, MD, Division of Cardiology, Mount Sinai Hospital Service, City Hospital Center, Elmhurst, NY 11373.

monary artery banding (31), ligation of a PDA (24) systemic to pulmonary anastomoses (18), resection of aortic coarctation (8) and atrial septectomy (8) Survival rate of all surgical procedures was 71 per cent At the end of 1 year of follow up, 139 patients were alive (54 per cent) However if patients who died without the opportunity for surgery or catheterization are excluded, survival rate was increased to 66 per cent

The authors wish to thank Drs Jay Levy and Marion Ledbetter of Madison Wisc Dr A C V Elston of La Crosse Wisc and Dr George Griese of Marshfield Wisc for their help and cooperation in obtaining data

REFERENCES

- 1 Fyler D C Cardiac catheterization cardiac surgery and the newborn infant—1969 *Pediatrics* 44:1 1969
- 2 Varghese P J Celermajer J Izukawa T Haller A J and Rowe R D Cardiac catheterization in the newborn Experience with 100 cases *Pediatrics* 44:24 1969
- 3 Waterston D H Treatment of Fallot's tetralogy in children under one year of age *Rozhl Chir* 44:181 1962
- 4 Richards M R Merritt K K Samuels M H and Langmann A G Congenital malformations of the cardiovascular system in a series of 6 053 infants *Pediatrics* 15:12 1955
- 5 Feldt R H Avasthey P Yoshimasu F Kurland K T and Titus J L Incidence of congenital heart disease in children born to residents of Olmsted County Minnesota 1950-1969 *Mayo Clin Proc* 46:794 1971
- 6 Coleman E H Serious congenital heart disease in infancy *Br Heart J* 27:42 1965
- 7 Mehri A Hirsch M S and Taussig H B Congenital heart disease in the neonatal period *J Pediatr* 65:1721 1964
- 8 Fyler D C Parisi L and Berman M New England Regional Infant Cardiac Program Abstracts of the 21st Annual Scientific Session American College of Cardiology *Am J Cardiol* p 264 1972
- 9 Stark J Hucin B Aberdeen E and Waterston D H Cardiac surgery in the first year of life Experience with 1049 operations *Surgery* 69:483 1971
- 10 Barratt Boyes B G Simpson M and Neutze J M Intracardiac surgery in neonates and infants using deep hypothermia with surface cooling and limited cardiopulmonary bypass *Circulation* 43 and 44 (Suppl 1) 125 1971
- 11 Cambell M Natural history of cyanotic malformations and comparison of all common cardiac malformations *Br Heart J* 34:3 1972

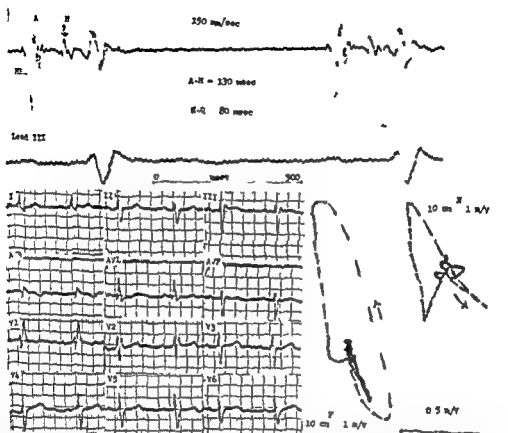


Fig 1 ECG VCG and His bundle electrogram (HBE) recorded from a patient in Group A 1. The initial forces are directed inferiorly and anteriorly with counterclockwise (CCW) rotation in the horizontal plane. The HBE demonstrates delayed conduction distal to the His deflection. HBE 1 = atrial depolarization. H = His bundle deflection. VCG H = horizontal plane. F = frontal plane.

designated Group A while those with a clockwise loop were designated Group B. There were 46 Group A and 19 Group B patients. The groups were then subdivided according to the initial vector in both the frontal and horizontal plane. A summary of significant findings is shown in Table 1.

Group A 1 (Fig 1) Initial forces anterior and inferior. Counterclockwise loop in the horizontal plane. There were 26 patients in this group: 20 men and six women with an average age of 69 years. The most significant findings were the high incidence of enlarged heart and hypertension (7/26) and the high incidence of syncope (12/26). First degree AV block was noted in four and transient complete heart block was noted in two. Nine of 11 His bundle recordings showed a prolonged H-Q interval (normal H-Q interval 35 to 55 msec). Each

of the patients with a prolonged H-Q interval had a history of dizziness or syncope.

Group A 2 (Fig 2) Initial forces directed superiorly in the frontal plane. Counterclockwise horizontal loop with initial force directed anteriorly. There were 11 patients in this group: five men and six women with an average age of 67 years. Hypertension was found in four and dizziness or syncope in five. There were two instances of first degree AV block and one of complete heart block. Three of six His bundle recordings showed a prolonged H-Q interval and each of these patients had a history of dizziness or syncope.

Group A 3 (Fig 3) Initial forces directed posteriorly and counterclockwise loop in the horizontal plane. There were nine patients in this group: seven men and two women with an average age of 70 years. The most

Table 3 Summary of significant findings in 65 patients

| Group | A1 | A2 | A3 | B1 | B2 | B3 |
|----------------------------------|----|----|----|----|----|----|
| Average age | 69 | 67 | 70 | 74 | 79 | 73 |
| Men | 20 | 5 | 7 | 6 | 4 | 2 |
| Women | 6 | 6 | 2 | 4 | 1 | 2 |
| Myocardial infarction (clinical) | 3 | 1 | 8 | 2 | 1 | 1 |
| Angina pectoris | 3 | 1 | 0 | 1 | 0 | 1 |
| Diabetes mellitus | 4 | 2 | 1 | 2 | 2 | 1 |
| Hypertension | 7 | 4 | 2 | 1 | 2 | 1 |
| Enlarged heart (radiographic) | 7 | 3 | 6 | 7 | 3 | 2 |
| Congestive heart failure | 4 | 3 | 6 | 5 | 3 | 1 |
| Dizziness/syncope | 12 | 5 | 2 | 3 | 1 | 0 |
| 3° A V block | 2 | 1 | 0 | 2 | 1 | 0 |
| 2° A V block | 0 | 0 | 1 | 0 | 0 | 0 |
| 1° A V block | 4 | 2 | 2 | 4 | 0 | 0 |
| H Q > 55 | 9 | 3 | 0 | 1 | 1 | 0 |
| A H > 140 | 1 | 1 | 1 | 0 | 0 | 0 |
| Total His recordings | 11 | 6 | 7 | 3 | 1 | 0 |

were greater than 160/90. If the patient was on anti hypertensive medication this information was obtained from the patient's previous record. Cardiomegaly was diagnosed radiographically if the cardiothoracic ratio was greater than 0.5. Dizziness and syncope have been grouped together. This symptom was recorded as being present only if non cardiac causes could be excluded. None of the patients had evidence of acute myocardial infarction. There were 44 men and 21 women. Their ages ranged between 31 and 93 years with an average age of 70 years.

The ECG criteria for inclusion in this study were as follows:

1. Total QRS duration of at least 0.12 seconds.

2. Mean frontal plane QRS axis of -30 degrees or greater. The majority of patients had a small initial R wave in Leads II, III and aVF. However this was not required as a criteria. Patients with QS complexes in these leads were included thus filling the criteria for left axis and not for left anterior hemiblock.

3. Evidence of right bundle branch block manifested by delay and slurring of the R wave in the right precordial leads and slurred S wave in the left precordial leads.

VCGs were recorded with the Frank lead system⁷ with patients in the supine position and the chest electrodes at the level of the fifth intercostal space. Frontal

horizontal, and left sagittal loops were recorded in addition to scalar X, Y, and Z. The VCGs were recorded on either a DR 12 Electronics for Medicine photographic recorder which displays a dash representing 4 msec or a Hewlett Packard oscilloscopic VCG with loops photographed with a Polaroid camera. In the latter instance each dash represented 2.5 msec. Conventional reference frames for the horizontal and frontal planes were used for angular measurements.

The following information was obtained from each projection of the QRS loop: (1) direction of inscription of the QRS loop, (2) orientation of the mean QRS forces, (3) direction of the initial 20 msec vector, and (4) direction and speed of inscription of terminal forces.

Twenty eight patients also had a His bundle electrogram which was recorded within one week of the VCG. His bundle studies were performed according to the method of Scherlag and associates.⁸ Paper speeds of 75 and 150 mm per second and a frequency range of 40 to 500 Hz were used.

Results

Patients were divided into two groups depending on the direction of rotation of the horizontal loop of the VCG. The terminology suggested by Salzman and colleagues⁵ was used. Patients with a counter clockwise loop in the horizontal plane were

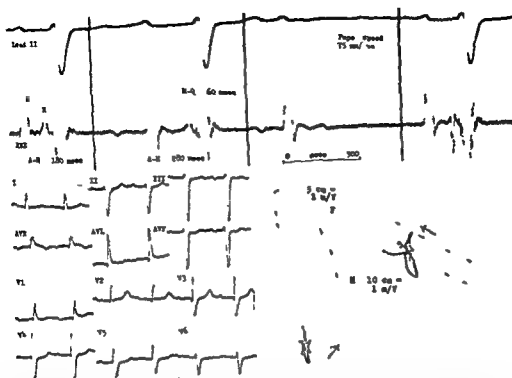


Fig 3 ECG, VCG and HBE recorded in a patient in Group A 3. The initial forces are directed inferiorly and posteriorly with a CCW loop in the horizontal plane. The HBE shows delay in conduction both proximal and distal to the His deflection (H). Mobitz Type I second degree $\Delta\Delta$ block is noted. See Fig 1 for explanation of abbreviations.

were slightly younger (average age 69 years) than the Group B patients (average age 75 years). The incidence of previous myocardial infarction (Group A 12/46 Group B 4/19) and hypertension (Group A 13/46 Group B 4/19) were similar. Diabetes mellitus was slightly more common in Group II (Group A 7/46 Group B 5/19). Cardiomegaly (Group A 16/46 Group B 12/19) and congestive heart failure (Group A 13/46 Group B 9/19) were more common in Group II. Episodes of syncope (Group A 19/46 Group B 4/19) and prolonged H-Q interval (≥ 5 msec or greater) (Group A 19/24 Group B 2/4) were more frequent in Group A patients. However in individual cases there was no definite relationship between prolonged H-Q interval and syncope.

Discussion

The combination of right bundle branch block and abnormal left axis deviation is

recognized in approximately 1 per cent of routinely interpreted ECGs.^{8,10} The most common etiology is coronary artery disease.^{11,12} Scanlon and colleagues¹³ found clinical evidence of coronary artery disease in 41 per cent of a group of patients with this electrocardiographic pattern. There is frequently obstruction of the left anterior descending coronary artery since the middle third of the right bundle and the anterior division of the left bundle are supplied by branches of this artery.¹³

Diffuse myocardial fibrosis involving the intraventricular conduction pathways is another etiologic factor described by Lenegre.¹ These patients are usually asymptomatic with no history of angina pectoris or valvular disease.¹

A small group of patients with this ECG pattern have calcific aortic stenosis and left ventricular hypertrophy. It is thought that the bifascicular block is probably due to a sclerotic degenerative process de-

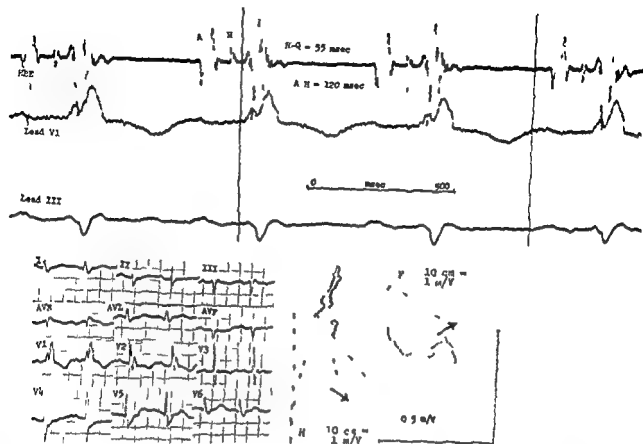


Fig 2 ECG VCG and HBE recorded from a patient in Group A 2. The initial forces are directed superiorly and anteriorly with a CCW loop in the horizontal plane. The HBE shows the conduction distal to the His deflection (H) to be at the upper limits of normal. See Fig 1 for explanation of abbreviations.

striking finding in this group was the high incidence of clinical myocardial infarction (8/9). There was also a high incidence of enlarged heart and congestive heart failure (6/9). There were two patients with dizziness or syncope; two patients had first degree A-V block. Six of seven His bundle recordings showed prolongation of the H-Q interval, but only one of these patients had a history of dizziness.

Group B 1 (Fig 4) Initial vectors directed inferiorly, anteriorly, and to the right. Clockwise loop in the horizontal plane. There were ten patients in this group: six men and four women with an average age of 74 years. The most striking findings were the high incidence of an enlarged heart (7/10) and congestive heart failure (5/10). Three patients had dizziness or syncope; four had first degree A-V block, and two had episodes of complete heart block. Only one of the three His bundle recordings showed evidence of H-Q prolongation. This patient did not have a history of dizziness or syncope.

Group B 2 (Fig 5) Initial forces directed superiorly in the frontal plane, clockwise loop in the horizontal plane. There were five patients in this group: four men and one woman with an average age of 79 years. Three out of five patients had both enlarged heart and congestive heart failure. One patient had dizziness or syncope. The one His bundle recording performed in this group showed a prolonged H-Q interval and was done on a patient with a history of dizziness.

Group B 3 (Fig 6) Initial forces directed posteriorly in the horizontal plane with a clockwise rotation. There were four patients in this group: two men and two women with an average age of 73 years. Two of these patients had enlarged heart. There were no instances of dizziness or syncope. No His bundle electrograms were recorded in this group.

Summary of results

A summary of results in the two basic groups is as follows. The Group patients

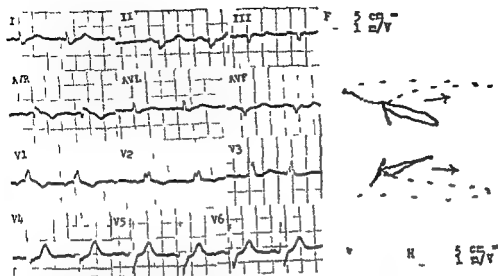


Fig 5 ECG and VCG recorded from a patient in Group B 2. The initial forces are directed superiorly and anteriorly with a CW initial loop in the horizontal plane. See Fig 1 for explanation of abbreviations.

tions found in order of decreasing frequency were syncope, myocardial infarction, hypertension, diabetes mellitus, and angina pectoris. A frequent occurrence of hypertension and diabetes mellitus was noted in the series of Scanlon and co-workers.¹¹

Since the VCG morphology varies, it was thought that various patterns might have some relationship to the clinical course. Salzman and colleagues⁵ reported that patients with a clockwise horizontal loop had a higher frequency of overt and symptomatic coronary disease and had a greater frequency of cardiomegaly, congestive heart failure, and death when compared to patients with a counterclockwise horizontal loop. Kulbertus and associates⁴ was unable to find any significant clinical difference between two similar groups of patients.

Salzman and colleagues⁵ suggested that true posterior wall infarction and fibrosis was an explanation of the anteriorly directed clockwise horizontal loop. Hugenholz and co-workers¹⁰ described four out of 33 cases with posterior basal myocardial infarction with a clockwise loop in the horizontal plane. Walsh and associates²¹ also described occasionally finding clockwise loops in the horizontal plane with posterior wall myocardial infarction.

An experimental study by Kulbertus

and colleagues⁴ showed that an identical clockwise loop in the horizontal plane could be produced in normal controls by stimulation of the posterior surface of the heart. A study by Cohen and co-workers²² also showed that this type of pattern may be produced by the introduction of atrial premature beats. Both clockwise and counterclockwise loops were produced. The presence or absence of underlying heart disease did not appear to affect the type of VCG pattern produced.

Several clinical differences were observed in our patients. Group A patients were slightly younger than Group B. Diabetes mellitus, cardiomegaly, and congestive heart failure were more common in Group B. The amount of cardiomegaly and congestive heart failure in Group B could not be explained by the incidence of hypertension and was thought to be due to more extensive myocardial damage. Episodes of syncope and prolonged H Q interval (55 msec or greater) were more frequent in Group A patients. However, in individual cases there was no definite relationship between prolonged H Q interval and syncope (Table II).

It is concluded that the ECG pattern of right bundle branch block with abnormal left axis deviation represents a wide spectrum of disease and anatomical defects.

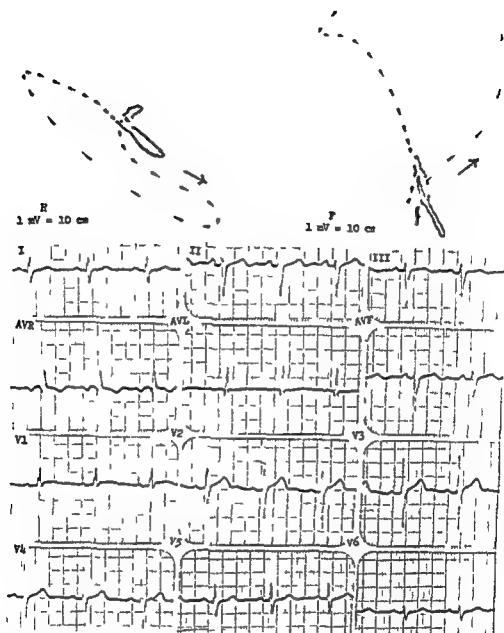


Fig 4 ECG and VCG recorded from a patient in Group II 1. The initial forces are directed inferiorly and anteriorly with a clockwise (CW) loop in the horizontal plane. See Fig 1 for explanation of abbreviations.

scribed by Lev¹⁵ which extends down from the aorta to involve the membranous septum and the top of the muscular septum.

Cardiomyopathy may also produce this pattern as seen in the case described by Castellanos and co workers¹⁶. Autopsy on this patient showed normal coronary arteries but wide spread areas of patchy and diffuse fibrosis. This ECG pattern may also be encountered following tricuspid valve replacement¹⁷ and repair of a ventricular septal defect¹⁸.

An important reason for recognizing this pattern is in predicting clinical complication such as complete heart block. Lasser Haft and Friedberg¹⁰ report an incidence of 10 per cent while Scanlon, Pryor and Blount¹² reported an incidence of 14 per cent. In a smaller series reported by Kulbertus and Collignon¹⁹ all 11 patients developed transient or permanent complete heart block and episodes of syncope. In 63 patients with this ECG pattern reported by Watt and Pruitt⁹ associated clinical condi-

Table II Comparison of clinical differences in patients in Groups A and B

| No | Age | Group | AH | HQ | A V Conduction | Dizziness/syncope |
|----|-----|-------|-----|-----|--|-------------------|
| 1 | 52 | A1 | 90 | 50 | | + |
| 2 | 83 | A1 | 130 | 35 | | |
| 3 | 84 | A1 | 110 | 70 | | + |
| 4 | 72 | A1 | 270 | 70 | 1 AV block | + |
| 5 | 76 | A1 | 140 | 80 | Intermittent CHB | + |
| 6 | 90 | A1 | 130 | 80 | 1 AV block | + |
| 7 | 60 | A1 | 80 | 100 | 1 AV block | + |
| 8 | 78 | A1 | 100 | 90 | | + |
| 9 | 81 | A1 | 110 | 60 | | + |
| 10 | 71 | A1 | 90 | 70 | CHB | + |
| 11 | 87 | A1 | 80 | 60 | | + |
| 12 | 85 | A2 | 120 | 60 | | + |
| 13 | 72 | A2 | 120 | 45 | Intermittent 1 AV block | |
| 14 | 40 | A2 | 110 | 50 | | + |
| 15 | 77 | A2 | 90 | 65 | CHB | + |
| 16 | 89 | A2 | 80 | 45 | | + |
| 17 | 78 | A2 | 90 | 60 | | + |
| 18 | 80 | A3 | 100 | 60 | | |
| 19 | 46 | A3 | 110 | 40 | | |
| 20 | 51 | A3 | 110 | 55 | | |
| 21 | 78 | A3 | 170 | 55 | 1 AV block and intermittent 2 AV block proximal to His potential | |
| 22 | 83 | A3 | 90 | 90 | | |
| 23 | 56 | A3 | 90 | 60 | | |
| 24 | 93 | A3 | AF | 70 | | |
| 25 | 70 | B1 | 100 | 50 | Intermittent 1 AV block | + |
| 26 | 75 | B1 | 120 | 45 | | + |
| 27 | 73 | B1 | 80 | 70 | | |
| 28 | 83 | B2 | 90 | 100 | CHB | + |

P is not on digitalis.

and hypertension (Group A 13/46 Group B 4/19) were similar. Diabetes mellitus was slightly more common in Group B (Group A 7/46 Group B 5/19). Cardiomegaly (Group A 16/46 Group B 12/19) and congestive heart failure (Group A 13/46 Group B 9/19) were more common in Group B. Episodes of syncope (Group A 19/46 Group B 4/19) and prolonged H Q interval (55 msec or greater) (Group A 19/24 Group B 2/4) were more frequent in Group A patients. However in individual cases there was no definite relationship between prolonged H Q interval and syncope.

It is concluded that this pattern represents a wide spectrum of disease and anatomical defects. The relationship of VCG morphology to clinical course does not appear to be close enough to predict the course of an individual patient or aid in clinical management.

The authors wish to thank Richard P. Lasser M.D. for his helpful suggestions in the course of this study.

REFERENCES

1. Wilson F N, Johnston F D and Barker P S. Electrocardiograms of an unusual type in right bundle branch block. *AM HEART J* 9:472 1934.
2. Lenegre J. Etiology and pathology of bilateral bundle branch block in relation to complete heart block. *Progr Cardiovasc Dis* 6:409 1964.
3. Unger P N, Lesser M E, Kugel V H and Lev M. The concept of masquerading bundle branch block. An electrocardiographic pathologic correlation. *Circulation* 17:397 1968.
4. Watt T B Jr, Freud G E, Durrer D and Pruitt R D. Left anterior arborization block combined with right bundle branch block in canine and primate hearts. An electrocardiographic study. *Circ Res* 22:57 1968.
5. Salzman T, Linn H and Pick A. Right bundle branch block with left axis deviation. *Br Heart J* 28:703 1966.

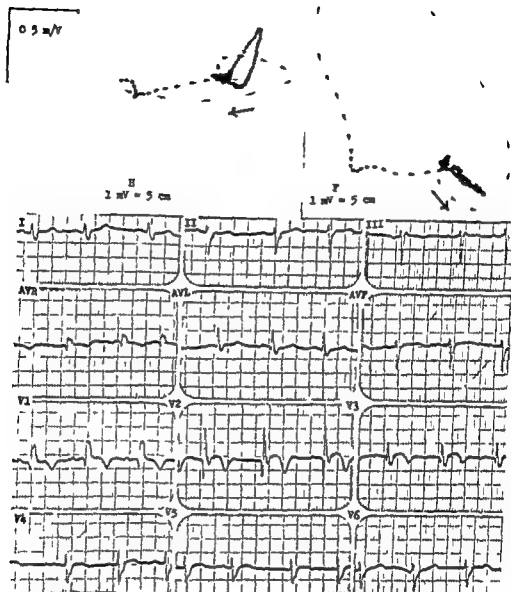


Fig 6 ECG and VCG recorded from a patient in Group B 3. The initial forces are directed inferiorly and posteriorly with a CW loop in the horizontal plane. See Fig 1 for explanation of abbreviations.

The abnormal initial forces in the frontal plane probably indicate inferior wall myocardial infarction which¹¹ coupled with the conduction defect suggests disease of both the right and left coronary arteries. The abnormal initial forces in the horizontal plane may indicate anterior septal myocardial infarction due to occlusion of the proximal left anterior descending coronary artery or they may be due to the left anterior hemiblock alone.²¹ The relationship of VCG morphology to clinical course does not appear to be close enough to predict the course of an individual patient or aid in clinical management.

Summary

Sixty five patients with a VCG pattern of complete right bundle branch block and left axis deviation (> -30 degrees) were studied. There were 46 patients with counterclockwise rotation (Group A) and 19 patients with clockwise rotation (Group B) in the horizontal plane. His bundle electrograms were recorded in 24 Group A patients and in four Group B patients.

The Group A patients were slightly younger (average age 69 years) than the Group B patients (average age 75 years). The incidence of previous myocardial infarction (Group A 12/46 Gr -4/19)

Left ventricular function in systole and diastole in constrictive pericarditis

B S Leus MB BCh

M S Gotsman MD FRCP FACC

Durban South Africa

Left ventricular (LV) function may be normal or depressed in constrictive pericarditis. Previous studies made in small groups of patients suggest that the mechanical function of the LV is abnormal while there is little information about diastole.¹⁻⁴

We have had an opportunity to study a large number of patients with constrictive pericarditis. This report will describe the function of the left ventricle in systole and diastole in a consecutive group of 30 patients.

Patients

We selected for study 30 consecutive patients with constrictive pericarditis (CP) who were evaluated before operation. Patients with atrial fibrillation were excluded as the varying R-R interval complicates analysis. One patient had atrial flutter with 4:1 atrioventricular block and a regular ventricular response.

The clinical, electrocardiographic and radiological features of the patients were typical and have been described in detail.⁴⁻¹¹ The patients were in cardiac failure and the important clinical features are summarized in Table 1. Twenty-seven patients had underlying tuberculous disease, one had amoebic infection of the pericardium and

in two constriction followed a penetrating stab wound of the chest. All the patients were treated with digitalis and diuretics and were fully digitalized at the time of study.

The operation of pericardiectomy was made in 18 patients; the remainder are presently receiving preoperative anti-tuberculous therapy or are being treated conservatively.

Methods

Data acquisition. Premedication with 10 mg diazepam (Valium) and 50 mg Pethidine was given one hour before study. Routine cardiac catheterization was performed through a percutaneous puncture of the right femoral artery and vein or a cutdown in the right antecubital fossa. Intravascular and intracardiac pressures were measured through Statham P23 Db bonded strain gauges using the mid-chest level as the zero reference. Pulsus paradoxus was measured in the ascending aorta during deep respiration. Pressures for calculation were the mean values recorded during several respiratory cycles. LV dp/dt was derived from LV pressure using an electronic analog differentiating amplifier without phase lag or distortion.

Cardiac output was measured by the

From the Cardiology Department and the University of Natal, Durban, South Africa.
Supported by a grant from the Medical Research Council of South Africa.
Received for publication Aug 28, 1972.
Reprint requests to Prof M S Gotsman, MRC, Durban Hospital, P B Jacobs, Natal, South Africa.

- 6 Kulbertus H Collignon P and Humblet N Vectorcardiographic study of QRS loop in patients with left superior axis deviation and right bundle branch block *Br Heart J* 38:386 1970
- 7 Frank E Accurate clinically practical system for spatial vectorcardiography *Circulation* 13:737 1956
- 8 Scherlag H J Law S H Helfant R H Berkowitz W D Stein E and Damato A N Catheter technique for recording His bundle activity in man *Circulation* 39:13 1969
- 9 Watt T B Jr and Pruitt R B Character cause and consequence of combined left axis deviation and right bundle branch block in human electrocardiograms *AM HEART J* 77:460 1969
- 10 Lasser R P Haft J I and Friedberg C K Relationship of right bundle branch block and marked left axis deviation (with left parietal or peri infarction block) to complete heart block and syncope *Circulation* 37 429 1968
- 11 Rosenbaum M B Types of right bundle branch block and their clinical significance *J Electrocardiography* 1 (Suppl 2):221 1968
- 12 McClenahan J B Significance of pronounced left axis deviation in presence of right bundle branch block *Calif Med* 110:378 1969
- 13 Scanlon P J Pryor R and Blount S G Right bundle branch block associated with left superior or inferior intraventricular block, *Circulation* 42:1123 1970
- 14 James T N The coronary circulation and conduction system in acute myocardial infarction *Progr Cardiovasc Dis* 10 410 1968
- 15 Lev M Anatomic basis for atrioventricular block *Am J Med* 37:742 1964
- 16 Castellanos A Jr Lemberg L Ioannides G and Salhanick N The vectorcardiogram in right bundle branch block co-existing with left ventricular focal block *Am J Cardiol* 18:705 1966
- 17 Aravindakshan V Elizarı M V and Rosenbaum M B Right bundle branch block in left anterior fascicular block (left anterior hemiblock) following tricuspid valve replacement *Circulation* 42:895 1970
- 18 Kulbertus A G Coyne J J Hallidie Smith K A Conduction disturbances before and after surgical closure of ventricular septal defect *AM HEART J* 77:123 1969
- 19 Kulbertus H and Collignon P Association of right bundle branch block with left superior or inferior intraventricular block *Br Heart J* 31 435 1969
- 20 Hugenholtz P G Forkner C E and Levine H D A clinical appraisal of the vectorcardiogram in myocardial infarction II The Frank system *Circulation* 23:825 1961
- 21 Walsh T J Tongson P M Stoddard E A and Massie E The vectorcardiographic QRS E loop findings in inferior posterior myocardial infarction *AM HEART J* 63:516 1962
- 22 Cohen S I Law S H Stein E Young M W and Damato A N Variations of aberrant ventricular conduction in man Evidence of isolated and combined block within the specialized conduction system *Circulation* 38 899 1968
- 23 Benchimol A and Desser K B Co-existing left anterior hemiblock and inferior wall myocardial infarction—vectorcardiographic features *Am J Cardiol* 29:7 1972
- 24 McHenry P I Phillips J F Fisch C and Corya B R Right precordial qrs pattern due to left anterior hemiblock *AM HEART J* 81:498 1971

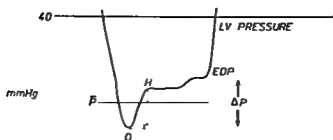


Fig 1 Ventricular diastolic pressure pulse tracing in a patient with constrictive pericarditis O is the lowest diastolic pressure and EDP is the instant of onset of left ventricular pressure rise H is the diastasis plateau and ΔP is the diastolic pulse pressure \bar{P} is the mean diastolic pressure where $\bar{P} = \frac{EDP - O}{2}$ mm Hg

correction made in the measurements. The systolic time intervals have two components: pre-ejection phase (PEP) from the earliest Q or R wave of the ECG to the instant of aortic pressure rise and left ventricular ejection time (LVET) from aortic valve opening to its closure. Total electromechanical systole ($Q S_2$) is the time from the onset of ventricular activation on the ECG until aortic valve closure.^{12, 13} The PEP/LVET ratio was also calculated; this ratio correlates with the left ventricular ejection fraction.¹⁶ PEP and LVET are heart rate dependent; we calculated ΔPEP and $\Delta LVET$ where the Δ value is the deviation from the normal regression lines of Weissler, Harris and Schoenfeld¹³ at a given heart rate. Our patients were digitalized and it is possible that we should have used the revised regression of Weissler and Schoenfeld¹⁷ allowing for digitalization. There is a beat to beat variation of time interval measurements in pulsus paradoxus and we used in our calculations the mean value recorded during several respiratory cycles.¹⁸

Ventricular volumes were measured from the uniplane cineangiogram using the method of Greene and associates¹⁹ and allowing for magnification. End-diastolic (EDV) and end-systolic (ESV) and angiographic stroke index (angio SI) were calculated and were normalized for body surface area. The ejection fraction and filling fraction were calculated where

$$\text{Ejection fraction (EF) (\%)} = \frac{EDV - ESV}{EDV} \times 100$$

and

$$\text{Filling fraction (FF) (\%)} = \frac{EDV - ESV}{ESV} \times 100$$

Left ventricular ejection rate was calculated where

$$\text{Ejection rate (ml/sec/1.73)} = \frac{\text{Angio SI (ml/beat/1.73)}}{LVET \text{ (msec)}} \times 10^3$$

$\Delta P/\Delta V$ was calculated where ΔP was divided by angiographic $SI \Delta V/\Delta P$ is the over all compliance of the ventricle in diastole.

The following derived measurements were calculated: Circumferential fiber shortening (C_f) where

$$C_f (\%) = \frac{C_{dss} \text{ (cm)} - C_{s2} \text{ (cm)}}{C_d \text{ (cm)}} \times 100$$

Mean velocity of circumferential fiber shortening (V_c) where

$$V_c \text{ (circ/sec)} = \frac{C_d \text{ (cm)} - C_{s2} \text{ (cm)}}{C_{dss} \text{ (cm)} \times LVET \text{ (msec)}} \times 10$$

and

$$C_d = \text{Inner equatorial LV diameter in diastole}$$

and

$$C_{s2} = \text{Inner equatorial LV diameter in systole}$$

Table I Clinical data

| Patient No | Initials | Age (yr) | Sex | Race | Etiology | JVP* (cm) | Hepato megaly (cm) | Edema | Ascites | Previous pericardial effusion |
|------------|----------|----------|-----|------|------------|-----------|--------------------|-------|---------|-----------------------------------|
| 1 | I M | 26 | M | W | TB | 8 | 8 | + | + | - |
| 2 | M W | 48 | M | B | TB | M | 4 | - | + | - |
| 3 | N N | 35 | M | B | Trauma | 10 | 8 | + | + | - |
| 4 | E N | 27 | M | B | TB | M | 5 | + | + | + |
| 5 | A M | 28 | F | B | TB | M | 12 | + | + | - |
| 6 | J M | 54 | M | B | TB | M | 6 | + | + | - |
| 7 | A S | 37 | M | W | TB | 10 | 3 | + | - | - |
| 8 | M N | 17 | M | B | TB | M | 8 | - | - | Previous pericardiectomy (12 yrs) |
| 9 | M N | 29 | M | B | TB | M | 4 | + | + | + |
| 10 | E N | 42 | M | B | TB | M | 6 | + | + | - |
| 11 | S N | 45 | M | B | TB | 4 | 4 | + | + | - |
| 12 | R E | 33 | M | W | TB | M | 8 | + | - | - |
| 13 | C M | 20 | F | B | TB | M | 4 | + | + | - |
| 14 | V N | 23 | F | B | TB | M | 6 | + | + | - |
| 15 | M M | 35 | M | B | Amoebiasis | M | 5 | + | + | + |
| 16 | J S | 23 | M | B | TB | 10 | 3 | + | + | + |
| 17 | G M | 49 | M | B | TB | 3 | 7 | - | + | - |
| 18 | R K | 31 | M | C | Trauma | 6 | - | - | - | + |
| 19 | S D | 50 | M | B | TB | 8 | 10 | + | + | - |
| 20 | M S | 30 | M | B | TB | M | 10 | + | + | - |
| 21 | M S | 55 | M | B | TB | 8 | 6 | + | + | - |
| 22 | J M | 50 | M | B | TB | M | 3 | + | + | - |
| 23 | A D | 48 | M | B | TB | 5 | 6 | + | + | - |
| 24 | B N | 28 | M | B | TB | M | 8 | + | + | - |
| 25 | M M | 14 | M | B | TB | 10 | 5 | + | + | + |
| 26 | K S | 27 | M | B | TB | 8 | 8 | - | - | + |
| 27 | S N | 27 | M | B | TB | 6 | 5 | + | + | - |
| 28 | S M | 44 | M | B | TB | M | 8 | + | + | - |
| 29 | J N | 43 | M | B | TB | M | 8 | + | + | + |
| 30 | G M | 46 | M | B | TB | M | 6 | + | + | - |

*Abbreviations B = Bantu C = Colored JVP = jugular venous pressure M = jugular venous pressure above angle of mandible in erect position TB = tuberculosis W = white

direct Fick method Routine cardiac catheterization was followed by left ventricular cineangiography in the right anterior oblique (RAO) view using a slow injection of 50 ml of 76 per cent Urografin

Definitions and data analysis The analysis of diastole is shown in Fig 1 *O* was the lowest diastolic pressure recorded but we cannot exclude artefact from mechanical overshoot with certainty The *H* point was the instant of pressure inflexion at the end of the rapid filling phase of the left ventricle and was clearly defined in all but three patients who had a rapid heart rate at the time of study and had insufficient time for diastasis End diastolic pressure (*LVEDP*) was the instant of onset of the rapid rise of ventricular pressure The mean rate of

pressure rise during the rapid filling phase was calculated by dividing the *OH* pressure difference by the *OH* time interval and was expressed in mm Hg per second ΔP was the pressure difference between *O* and *LVEDP* Mean diastolic pressure (\bar{P})

$$\text{was } \frac{LVEDP - O}{2} \text{ mm Hg}$$

The time intervals of the cardiac cycle were measured from the electrocardiogram phonocardiogram and from a simultaneous pressure recording in the ascending aorta The delay time of the catheter manometer system was determined from a comparison of the dicrotic notch of the aortic pressure tracing with the aortic component of the second heart sound and the appropriate

Table II Hemodynamic and angiographic measurements

| Patient No | Pressures (mm Hg) | | | | | | | | | | Mean D.P.R. (mm Hg/sec) | Peak LV dp/dt (mm Hg/sec) | A-V O ₂ diff (ml/L) |
|--------------|-------------------|------------------|---------|----------|----------|-----------|------------|----------|-----------|------------|----------------------------|------------------------------|-----------------------------------|
| | MAP* | Pulsus paradoxus | Mean RA | RV LVEDP | Mean PAP | Mean PCWP | LV O point | LV LVEDP | \bar{P} | ΔP | | | |
| 1 | 92 | 5 | 25 | 28 | 36 | 28 | 10 | 30 | 20 | 20 | 188 | 1 117 | 40 |
| 2 | 94 | 11 | 18 | 26 | 35 | 28 | 20 | 33 | 27 | 13 | 90 | 1 480 | 44 |
| 3 | 80 | — | 11 | 14 | 19 | 11 | 8 | 16 | 12 | 8 | 85 | 1 150 | 37 |
| 4 | 86 | 13 | 23 | 22 | 33 | 23 | 22 | 32 | 27 | 10 | 57 | — | 41 |
| 5 | 82 | 14 | 18 | 20 | 28 | 21 | 16 | 23 | 20 | 7 | 138 | — | 23 |
| 6 | 106 | 30 | 26 | 28 | 31 | 26 | — | — | — | — | — | — | 33 |
| 7 | 88 | 12 | 22 | 28 | 33 | 26 | 8 | 20 | 17 | 6 | 110 | 1 060 | 37 |
| 8 | 88 | 12 | 23 | 30 | 36 | 26 | — | — | — | — | — | — | 37 |
| 9 | 104 | 10 | 6 | 6 | 20 | 8 | 4 | 15 | 10 | 11 | 58 | — | 47 |
| 10 | 80 | 10 | 11 | 20 | 40 | 17 | 12 | 20 | 16 | 8 | 63 | — | 35 |
| 11 | 90 | 10 | 7 | 10 | 20 | — | 8 | 13 | 11 | 5 | 27 | 906 | 41 |
| 12 | 76 | — | 22 | 25 | 33 | 25 | 9 | 24 | 17 | 15 | 144 | 1 050 | 40 |
| 13 | 95 | 24 | 19 | 18 | 28 | 26 | — | — | — | — | 13 | 2 114 | 31 |
| 14 | 101 | 14 | 21 | 27 | 27 | 21 | 15 | 26 | 21 | 11 | 151 | 1 255 | 31 |
| 15 | 92 | 20 | 8 | 15 | 17 | 9 | 7 | 17 | 12 | 10 | 86 | — | 34 |
| 16 | 105 | 28 | 23 | 30 | 35 | 27 | 18 | 27 | 23 | 9 | 63 | — | 31 |
| 17 | 100 | 15 | 1 | 3 | 10 | 4 | — | — | — | — | — | — | 31 |
| 18 | 93 | 10 | 11 | 12 | 20 | 12 | 10 | 14 | 10 | 4 | 60 | 1 208 | 27 |
| 19 | 85 | 10 | 21 | 22 | 30 | 20 | 12 | 19 | 16 | 7 | 50 | — | 43 |
| 20 | 105 | 55 | 19 | 19 | 34 | 25 | 12 | 25 | 19 | 13 | 144 | — | 41 |
| 21 | 87 | 10 | 8 | 8 | 19 | 16 | 7 | 16 | 10 | 9 | 40 | — | 39 |
| 22 | 80 | 20 | 18 | 10 | 26 | 7 | 8 | 19 | 14 | 11 | 69 | 1 070 | 38 |
| 23 | 100 | 24 | 18 | 20 | 25 | 19 | 14 | 24 | 19 | 10 | 143 | 1 478 | 35 |
| 24 | 75 | 15 | 20 | 20 | 30 | 25 | 25 | 35 | 30 | 10 | 167 | — | 60 |
| 25 | 87 | 31 | 17 | 20 | 30 | 21 | 11 | 21 | 16 | 10 | 125 | — | 37 |
| 26 | 110 | 35 | 20 | 20 | 30 | 20 | 19 | 30 | 25 | 11 | 100 | — | 54 |
| 27 | 90 | 25 | 16 | 20 | 20 | 15 | 15 | 22 | 19 | 7 | 50 | — | 45 |
| 28 | 90 | 20 | 16 | 23 | 22 | 22 | 15 | 24 | 20 | 9 | 67 | — | 47 |
| 29 | 105 | 60 | 24 | 30 | 33 | 30 | 11 | 28 | 21 | 14 | 214 | — | 85 |
| 30 | 85 | 18 | 27 | 21 | 33 | 28 | 20 | 27 | 21 | 7 | 60 | — | 51 |
| Mean | 91 | 21 | 18 | 20 | 28 | 20 | 13 | 23 | 18 | 10 | 100 | 1 314 | 41 |
| ±SD | 9 | 13 | 7 | 7 | 7 | 7 | 5 | 7 | 5 | 3 | 50 | 337 | 13 |
| Normal value | 90 ± 15 | — | 3 ± 2 | 4 ± 3 | 14 ± 5 | 9 ± 4 | 3 ± 2 | 9 ± 4 | 6 ± 3 | 6 ± 3 | — | 1 600 ± 400 | 41 ± 15 |
| Ref | 37 | — | 37 | 37 | 37 | 37 | 38 | 37 | 38 | 38 | — | 38 | 39 |

*Abbreviations: Angio SI = angiographic stroke index; A-V O₂ Diff = arterial-venous oxygen difference; Cf = circumferential fiber shortening (fraction); FF = filling fraction; LV dp/dt = 1st derivative of left ventricular pressure; LV LVEDP = left ventricular end diastolic pressure; change (LV LVEDP); PCWP = pulmonary capillary wedge pressure; PAP = pulmonary artery pressure; RA = right atrial pressure; velocity of circumferential fiber shortening.

Pericardial thickening obscured the outline of the left ventricle and true LV wall thickness and mid wall circumference could not be measured.

The LV stroke work index (SWI) was calculated where

$$\text{LV SWI (Gm M/beat/M}^2\text{)} = \frac{\text{Angio SI (ml/beat/M}^2\text{)} \times (\text{MAP} - \text{LV LVEDP}) (\text{mm Hg}) \times 0.0136}{\text{M}^2}$$

Statistical analysis The results were analyzed

using a Wang 700 programmable calculator and standard statistical methods.

Critique of methods Respiratory gymnastics and the subsequent delivery of a large volume of contrast medium during angiography disturbs the steady state of the circulation.²¹ Angiographic volumes are used as a standard of reference for measurement of ventricular volumes but we are hesitant to accept this without reservation. We used only technically perfect angio-

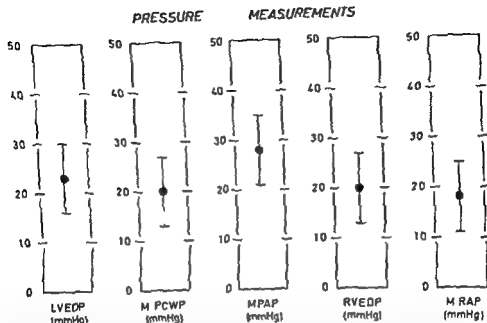


Fig 3 Pressure measurements in constrictive pericarditis. The ventricular end-diastolic pressures were elevated and were 2 to 3 mm Hg higher than the corresponding atrial pressures. Left heart pressures were 2 to 3 mm Hg higher than the corresponding right heart pressures. Mean pulmonary artery pressure (MPAP) was slightly elevated (28 ± 7 mm Hg). Mean values \pm 1 SD are shown.

striction and very small volumes had a reduced LV ejection rate (LVER) there was a significant linear correlation between LVER and EDV.

The linear relationship between EDV and angio SI was a reflection of a fairly constant ejection fraction (68 ± 9 per cent) which was maintained even at a low EDV. This relationship is shown in the upper panel of Fig 6.

Circumferential fiber shortening (C_1) and mean velocity of circumferential fiber shortening (V_1) were normal or slightly reduced ($C_1 = 28 \pm 9$ per cent $V_1 = 1.18 \pm 0.42$ circ/sec) (Fig 7). This was expected as they measure relative fiber shortening and correct for the reduced EDV; the measurements were related to ejection fraction and left ventricular ejection time which were within the range of normal.

Stroke work. Left ventricular stroke work index was reduced (mean LVSWI $= 29.2 \pm 13.8$ Gm Ml/beat/M²) it correlated with the severity of constriction as assessed by a reduction in EDV ($P < 0.001$) (Fig 8). Mean arterial pressure was normal and LVEDP was elevated but the fall in SWI reflected the decrease in SI.

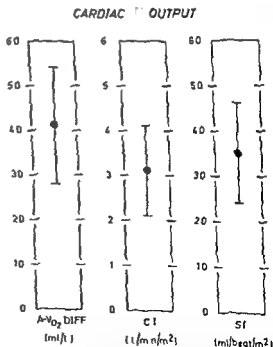


Fig 4 Measurements of mean arteriovenous oxygen difference (A-V O₂ diff), cardiac index (CI) and stroke index (SI) were normal but patients with severe constriction had a wide A-V O₂ difference and a low cardiac and stroke index. Mean values \pm 1 SD are shown.

LV DIASTOLIC PRESSURES

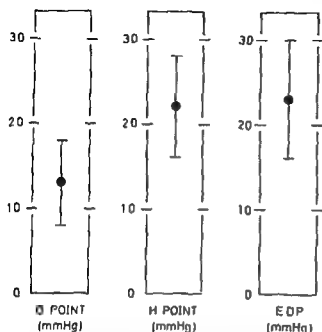


Fig 2 Left ventricular diastolic pressures in constrictive pericarditis (CP). The pressures were elevated with a dip and plateau configuration. Mean values \pm 1 SD are shown.

prominent feature, however all the patients had a high venous filling pressure with severe systemic congestion and fluid retention even though their disability grading was mild or moderate. Some patients were not severely disabled after prolonged bedrest and intensive diuretic therapy. It is not always possible in constrictive pericarditis to correlate abnormalities of ventricular function with shortness of breath.

Pressure measurements Figs 2 and 3 summarize the pressure measurements in patients with CP. Left ventricular O H and EDP were 13 ± 5 mm Hg, 22 ± 6 mm Hg, and 23 ± 7 mm Hg. They were elevated and abnormal and 2 to 3 mm higher than the corresponding pressures in the right ventricle (RVEDP = 20 ± 7 mm Hg). The left atrium and pulmonary vein were entered retrogradely in several patients; the two pressures were identical and were equal to the pulmonary capillary wedge pressure (PCWP) recorded at the same time and there was no evidence of constriction of the orifices of the pulmonary veins or of annular constriction of the mitral valve. Mean PCWP (20 ± 7 mm Hg) and mean right atrial (RA) pressure (18 ± 7 mm Hg) were identical to the cor-

responding pressure pulse contours in the two ventricles in diastole, but it is noteworthy that the mean atrial pressure was lower than the ventricular EDP. This reflects the importance of atrial systole in filling the ventricle and raising the EDP.

The mean pulmonary artery pressure (MPAP) was 28 ± 7 mm Hg and was only slightly elevated despite the elevation of LVEDP and PCWP. The absence of important pulmonary arterial hypertension contrasts these patients with those who have LV failure from coronary artery disease or congestive cardiomyopathy and in whom severe pulmonary hypertension accompanies a similar elevation of LVEDP.

The cardiac output The mean arteriovenous oxygen difference (A-V O₂ diff) was normal but there was a large scatter in individual patients (41 ± 13 ml/L) and several patients had a wide difference. Mean cardiac index (CI) and stroke index (SI) were at the lower limit of normal (CI = 3.1 ± 1.0 L/min/M²; SI = 35 ± 11 ml/beat/M²) in patients with severe constriction; they were low (Fig 4).

LV dp/dt Peak LV dp/dt was normal or slightly reduced ($1,314 \pm 337$ mm Hg/sec).

Angiographic volume measurements Left ventricular end diastolic (EDV) end systolic (ESV), and stroke volumes were reduced (EDV = 46 ± 22 ml/M²; ESV = 14 ± 9 ml/M²; and angio SI = 32 ± 16 ml/M²) (Fig 5). These volumes reflected the degree of constriction and the patients could be divided into three groups on arbitrary criteria: severe (EDV < 25 ml/M²), moderate (EDV 25 to 50 ml/M²), and mild (EDV > 50 ml/M²).

A significant linear relationship between EDV and angiographic SI is shown in Fig 6. Patients with severe constriction and a very small EDV had a corresponding decrease in stroke index (SI = 0.68 EDV + 0.86 ± 4.50 ; $r = +0.96$; $p < 0.001$). The fall in EDV and SI seem to be the most reliable indices of the severity of constriction. Patients with more severe constriction in general had a larger pulsus paradoxus; pulsus paradoxus was difficult to interpret since it depended on the amount of intrathoracic pressure change.

The mean left ventricular ejection rate was at the lower limit of normal (124 ± 63 ml/sec/M²) but patients with severe con-

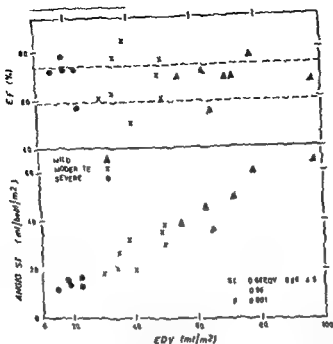


Fig 6 Linear relationship between end-diastolic volume (EDV) and angiographic stroke index (SI). The patients were divided into three arbitrary groups on the basis of their EDV: mild ($EDV > 30 \text{ ml/m}^2$), moderate ($25 \text{ to } 30 \text{ ml/m}^2$) and severe ($EDV < 25 \text{ ml/m}^2$). Patients with severe disease had a low EDV and a low SI. Ejection fraction (EF) was within the range of normal (dotted lines - upper panel).

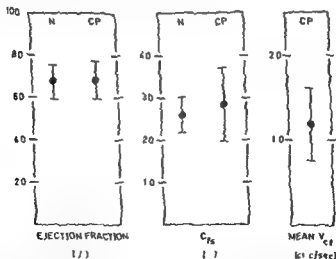


Fig 7 Angiographic measurements of fractional fiber shortening in normal subjects (N) and in patients with constrictive pericarditis (CP). Ejection fraction, circumferential fiber shortening (C_F) were normal in CP. Mean velocity of circumferential fiber shortening (V_c) was slightly reduced.

(\bar{P}) did not differentiate between mild and severe constriction.

The relationship between the rate of pressure rise in early diastole and the LVEDP is shown in Fig 16. The rate of

diastolic pressure rise was a function of the absolute ventricular pressure but was not related to EDV and the degree of constriction.

Filling fraction was normal. This mea-

LV VOLUMES

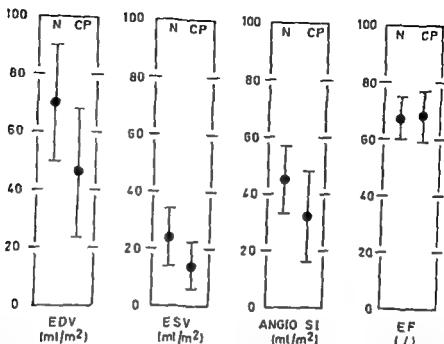


Fig 5 Angiographic left ventricular volume measurements in constrictive pericarditis (CP) and in normal subjects (N).¹⁰ End diastolic, end systolic and stroke volumes were reduced in CP but ejection fraction (EF) was normal.

Systolic time intervals PEP, LVET, and QS_2 were normal or marginally prolonged (Table III) and the relationships to heart rate are shown in Figs 9, 10, and 11. The figures show the regression of Weissler, Harris and Schoenfeld¹⁵ for normal subjects and of Weissler and Schoenfeld¹⁷ for normal digitalized subjects. The negative correlation between PEP and LV dp/dt is shown in Fig 12. There was no significant correlative relationship between LVET and SI but patients with a small stroke index always had a shortened LVET. The values of PEP/LVET clustered around the normal regression of Garrard, Weissler and Dodge¹⁸ since ejection fraction was normal.

Left ventricular compliance The LV diastolic pressure pulse showed an early diastolic dip (O point), a rapid filling wave ending at the H point, a significant plateau in diastasis and an important wave A. A detailed analysis of the diastolic pressures and the relationship to end diastolic volume (EDV) is shown in Fig 13. This shows that patients with severe constriction (EDV less than 25 ml/M²) had high diastolic pressures and a small diastolic pulse pressure (ΔP). Patients with mild constriction (EDV >50 ml/M²) also had

high end diastolic pressures but a wider diastolic pulse pressure (ΔP). Paradoxically patients with moderately severe constriction had less elevation of the diastolic pressures. ΔP was intermediate between the two groups. The reason for this discrepancy is not clear. We suggest that patients with moderate and severe constriction received more vigorous therapy than patients in the mild group; only patients with moderate disease were able to respond and reduce their blood volume and diastolic pressures.

The compliance (distensibility index) of the left ventriculo-pericardial system ($\Delta V/\Delta P$) was reduced and was related to EDV in a linear fashion (Fig 14). Patients with severe constriction had a small EDV and a low compliance.²⁰

Fig 15 shows the relationship between $\Delta P/\Delta V$ and \bar{P} to show the passive elastic modulus of the ventriculo-pericardial system measured according to the method of Drimond and Forrester and co-workers.^{21, 22} Patients with severe disease had a low compliance and a high passive elastic

$$\text{modulus} = \frac{(\Delta P/\Delta V)}{\bar{P}} \quad \text{Mean diastolic pressure}$$

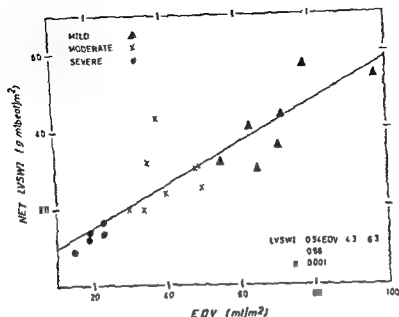


Fig 8 Linear relationship between net left ventricular stroke work index ($LVS\bar{W}$) and LV end-diastolic volume (EDV). Patients with severe constriction and a small EDV had a low net $LVS\bar{W}$.

by pericardiectomy.^{2,11,28} Ventricular filling pressures fall and cardiac output improves but they do not return to normal in the immediate postoperative period. The high venous filling pressure persists for 2 to 3 weeks and returns to normal in 2 to 3 months. Harrison, Crawford and Lau² made similar sequential studies of left ventricular function in a patient studied 7 weeks and 13 months after operation.

There are few reports which describe measurements of the mechanical function of the ventricle. Stroke output is limited and this has been attributed to a myocardial factor. Harvey and associates¹ in 2 patients decreased RV filling pressure by treatment with digitalis and diuretics—cardiac output increased in one and decreased in the other; the change was equated with an improvement in myocardial performance.¹ A simultaneous fall in blood volume occurred and is a more likely cause for the fall in pressure. This change is in keeping with our own clinical experience: digitalis and diuretic therapy produces a small but significant decrease in venous pressure, hepatic enlargement and edema, a non-specific consequence of a moderate decrease in blood volume.

Vogel and colleagues³ studied ventricular

dynamics and showed a significant reduction in EDV , ESV , SI , EF , C_r and V_{e1} . Their normal range of values for EF is high but comparative figures are valid and their data suggest that muscle fibers are compressed and that shortening is diminished in severe constriction.

Albers, Hugenholtz and Nadas¹¹ studied two patients with CP and atrial septal defect (ASD). The ejection fraction in one patient was 37 per cent before operation and returned to normal after pericardiectomy. The return to normal excludes an important myocardial factor and we believe that the ASD acted as a vent for the left atrium, decompressed the heart into the systemic circulation and permitted more severe constriction.

Dines, Edwards and Burchell¹⁰ studied 11 necropsy cases of CP and showed myocardial atrophy with a uniform decrease in diameter of myocardial fibers. Atrophy was not confined to the epicardial region adjacent to the constricting material. In CP intraventricular pressure and wall thickness are normal but cavity size is diminished, systolic wall stress is low, stroke output and stroke work are decreased and these factors may be responsible for disuse muscle fiber atrophy, analogous

Table III Time interval measurements

| Patient No | PEP | ΔPEP | ΔPEP _Δ | LVET | ΔLVET | ΔLVET _Δ | Q-S ₂ | ΔQ-S ₂ | ΔQ-S _{2Δ} | HR | PEP/LVET |
|--------------|--------|--------|-------------------|--------|--------|--------------------|------------------|-------------------|--------------------|-------------|----------|
| 1 | 150 | +53.0 | +68.0 | 300 | +31.5 | +46.5 | 450 | +82.5 | +112.5 | 85 | 0.50 |
| 2 | 90 | -7.8 | -7.2 | 280 | +8.1 | +23.1 | 370 | -1.7 | +28.3 | 83 | 0.32 |
| 3 | 120 | +22.2 | +37.2 | 200 | -71.9 | -56.9 | 320 | -51.7 | -21.7 | 83 | 0.60 |
| 4 | 90 | -3.4 | +11.6 | 250 | -3.2 | +11.8 | 340 | -8.6 | +21.4 | 94 | 0.35 |
| 5 | 90 | +9.0 | +24.0 | 260 | +50.0 | +65.0 | 350 | +61.0 | +91.0 | 130 | 0.35 |
| 6 | 80 | -9.8 | +5.2 | 210 | -27.9 | -12.9 | 290 | -39.7 | -9.7 | 103 | 0.38 |
| 7 | 80 | -25.8 | -10.8 | 260 | -25.9 | -10.9 | 360 | -53.7 | -23.7 | 63 | 0.29 |
| 8 | 110 | +12.2 | +27.2 | 260 | -11.9 | +3.1 | 370 | -1.7 | +28.3 | 83 | 0.49 |
| 9 | 70 | -25.8 | -10.8 | 290 | +28.6 | +41.6 | 360 | -1.2 | +28.8 | 88 | 0.74 |
| 10 | 90 | +1.8 | +16.8 | 240 | +8.9 | +23.9 | 330 | +5.7 | +38.7 | 107 | 0.38 |
| 11 | — | — | — | — | — | — | — | — | — | — | — |
| 12 | 70 | -29.4 | -14.4 | 260 | -18.7 | -3.7 | 330 | -50.1 | -20.1 | 79 | 0.27 |
| 13 | 50 | -31.0 | -16.0 | 220 | +10.0 | +25.0 | 270 | -19.0 | +11.0 | 130 | 0.23 |
| 14 | 83 | -7.2 | +7.8 | 250 | +3.2 | +16.2 | 333 | -2.0 | +28.0 | 107 | 0.33 |
| 15 | 120 | +9.0 | +44.0 | 240 | -3.0 | +12.0 | 360 | +24.0 | +54.0 | 100 | 0.50 |
| 16 | 140 | +46.6 | +61.6 | 230 | -23.2 | -8.2 | 370 | +21.4 | +51.4 | 94 | 0.61 |
| 17 | 110 | +12.2 | +27.2 | 280 | +8.1 | +23.1 | 390 | +18.3 | +48.3 | 83 | 0.39 |
| 18 | 90 | -1.0 | +14.0 | 240 | -3.0 | +12.0 | 330 | -6.0 | +24.0 | 100 | 0.31 |
| 19 | 70 | -19.8 | -4.8 | 260 | +22.1 | +37.1 | 330 | +0.3 | +30.3 | 103 | 0.27 |
| 20 | 80 | -14.6 | +0.4 | 260 | +1.7 | +16.7 | 340 | -14.9 | +15.1 | 91 | 0.31 |
| 21 | 130 | +34.2 | +49.2 | 250 | -13.4 | +1.6 | 380 | +18.8 | +48.8 | 88 | 0.52 |
| 22 | 80 | -21.8 | -6.8 | 290 | +1.1 | +16.1 | 370 | -27.7 | +7.3 | 73 | 0.79 |
| 23 | 90 | +9.0 | +24.0 | 210 | +9.5 | +24.5 | 300 | +16.5 | +46.5 | 125 | 0.43 |
| 24 | 140 | +45.4 | +60.4 | 240 | -18.3 | -3.3 | 380 | +25.1 | +55.1 | 91 | 0.58 |
| 25 | 70 | -13.0 | +2.0 | 230 | +21.0 | +36.0 | 300 | +6.0 | +36.0 | 120 | 0.30 |
| 26 | 90 | +5.0 | +20.0 | 200 | -17.5 | -2.5 | 290 | -14.5 | +15.5 | 115 | 0.45 |
| 27 | 120 | +25.4 | +40.4 | 260 | +1.7 | +16.7 | 380 | +25.1 | +55.1 | 91 | 0.46 |
| 28 | 110 | +4.2 | +19.2 | 300 | -5.9 | +9.1 | 410 | -3.7 | +26.3 | 63 | 0.37 |
| 29 | 120 | +31.8 | +46.8 | 210 | -21.1 | -6.1 | 330 | +8.7 | +38.7 | 107 | 0.57 |
| 30 | 50 | -44.6 | -29.6 | 280 | +21.7 | +36.7 | 330 | -24.9 | +5.1 | 91 | 0.18 |
| Mean | 95 | +3.0 | +18.0 | 251 | -1.4 | +13.6 | 347 | +0.0 | +30.0 | 96 | 0.39 |
| ± SD | 25 | 25.4 | 25.4 | 29 | 23.1 | 23.1 | 39 | 30.3 | 30.3 | 18 | 0.12 |
| Normal value | 0 ± 12 | 0 ± 11 | 0 ± 10 | 0 ± 10 | 0 ± 10 | 0 ± 10 | 0 ± 14 | 0 ± 14 | 0 ± 14 | 0.35 ± 0.04 | |

*Abbreviations: HR = heart rate (beats/min); LVET = left ventricular ejection time (msec); PEP = pre ejection phase (msec).
 Δ value = deviation from normal when corrected for heart rate¹². A value_Δ = deviation from normal when corrected for heart rate and digitalization¹⁷.

surement was related to ejection fraction

($EF = \frac{1}{1 - EF} - 1$) the calculation mag

nified small errors in systolic measurements

Discussion

Review of the literature There are several excellent reviews of the aberrations of pressure and flow in constrictive pericarditis^{1, 2, 10, 11, 27}. Cardiac and stroke output are low at rest and cannot increase normally on exercise, ventricular diastolic pressures are elevated with a dip and plateau configuration and the two ventricular diastolic pressure pulse contours are very similar. Important pulmonary hypertension does not

occur and this may be due to the inability of the right ventricle to increase its output on exercise.

Kesteloot and Denef¹³ described a syndrome of severe constriction of rapid onset and short duration associated with a high early diastolic ventricular pressure loss of the y descent and a hemodynamic pattern resembling myocardial fibrosis¹⁰. These patients have severe constriction and correspond to our patients with an EDV of less than 25 ml/M² and a narrow diastolic pressure pulse measurement of ventricular volumes clearly distinguishes these patients from those with primary myocardial disease.

The hemodynamic pattern is modified

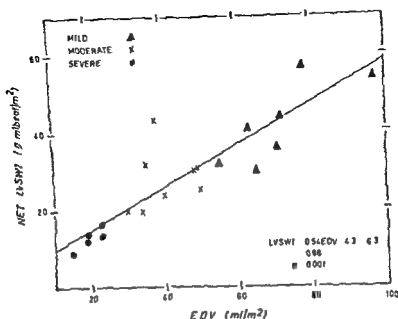


Fig. 8 Linear relationship between net left ventricular stroke work index (LVSWI) and LV end-diastolic volume (EDV). Patients with severe constriction and a small EDV had a low net LVSWI.

by pericardiectomy.^{2,11,28} Ventricular filling pressures fall and cardiac output improves but they do not return to normal in the immediate postoperative period. The high venous filling pressure persists for 2 to 3 weeks and returns to normal in 2 to 3 months. Harrison, Crawford and Lau² made similar sequential studies of left ventricular function in a patient studied 7 weeks and 13 months after operation.

There are few reports which describe measurements of the mechanical function of the ventricle. Stroke output is limited and this has been attributed to a myocardial factor. Harvey and associates¹ in 2 patients decreased RV filling pressure by treatment with digitalis and diuretics—cardiac output increased in one and decreased in the other; the change was equated with an improvement in myocardial performance.¹ A simultaneous fall in blood volume occurred and is a more likely cause for the fall in pressure. This change is in keeping with our own clinical experience: digitalis and diuretic therapy produces a small but significant decrease in venous pressure, hepatic enlargement and edema is non-specific consequence of a moderate decrease in blood volume.

Vogel and colleagues³ studied ventricular

dynamics and showed a significant reduction in EDV, ESV, SI, EF, C_1 and $V_{1/2}$. Their normal range of values for EF is high but comparative figures are valid and their data suggest that muscle fibers are compressed and that shortening is diminished in severe constriction.

Albers, Hugenholz and Nadas¹⁹ studied two patients with CP and atrial septal defect (ASD). The ejection fraction in one patient was 37 per cent before operation and returned to normal after pericardiectomy. The return to normal excludes an important myocardial factor and we believe that the ASD acted as a vent for the left atrium, decompressed the heart into the systemic circulation and permitted more severe constriction.

Dines, Edwards and Burchell¹⁰ studied 11 necropsy cases of CP and showed myocardial atrophy with a uniform decrease in diameter of myocardial fibers. Atrophy was not confined to the epicardial region adjacent to the constricting material. In CP intraventricular pressure and wall thickness are normal but cavity size is diminished, systolic wall stress is low, stroke output and stroke work are decreased and these factors may be responsible for disuse muscle fiber atrophy analogous

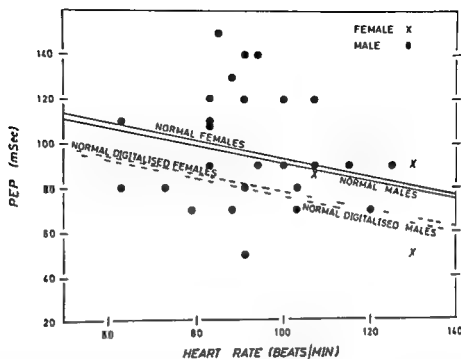


Fig 9 Relationship between pre ejection phase (PEP) and heart rate (HR) in patients with constrictive pericarditis. The regression lines of Weissler Harris and Schoenfeld¹⁶ for normal undigitalized subjects and of Weissler and Schoenfeld¹⁷ for normal digitalized subjects are shown. PEP was normal or slightly prolonged in CP.

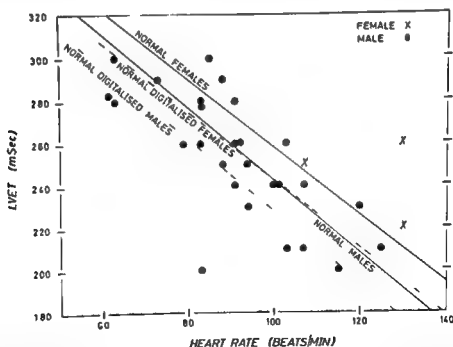
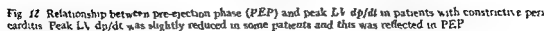


Fig 10 Relationship between left ventricular ejection time (LVET) and heart rate (HR). Regressions are shown for normal digitalized and undigitalized subjects.¹²⁻¹⁷ LVET was normal or slightly reduced in CP.

to skeletal muscle atrophy following immobilization of a limb.¹¹ This may account for the delayed return of the hemodynamics to normal following pericardiectomy when ventricular volumes, wall

stress and stroke work are returned to normal. Dines and colleagues¹⁰ selected a specific group of patients whose constriction was so severe that they died and were studied at autopsy.



Present study

SYSTOLIC FUNCTION The end-diastolic end systolic and stroke volumes of the left ventricle reflected the degree of constriction patients with severe disease had small volumes Patients with severe constriction and a low EDV had a low stroke index

stroke work index and left ventricular ejection rate

Ejection fraction was normal. In health and disease the ejection fraction is a sensitive index of ventricular dysfunction.²² Our results show that absolute fiber shortening is reduced and related to end diastolic

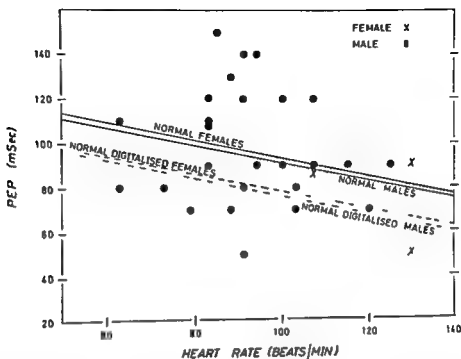


Fig 9 Relationship between pre ejection phase (PEP) and heart rate (HR) in patients with constrictive pericarditis. The regression lines of Weissler, Harris and Schoenfeld¹⁵ for normal undigitalized subjects and of Weissler and Schoenfeld¹⁷ for normal digitalized subjects are shown. PEP was normal or slightly prolonged in CP.

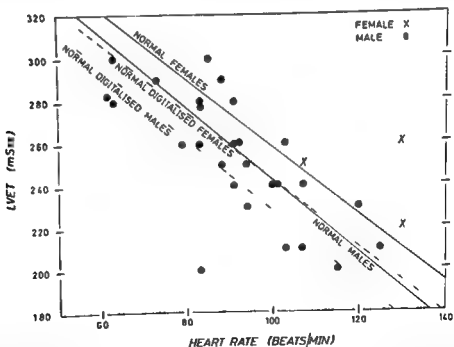
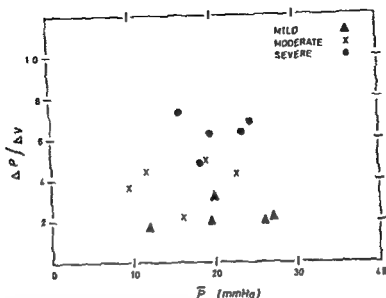


Fig 10 Relationship between left ventricular ejection time (LVET) and heart rate (HR). Regressions are shown for normal digitalized and undigitalized subjects.^{15, 17} LVET was normal or slightly reduced in CP.

to skeletal muscle atrophy following immobilization of a limb.³¹ This may account for the delayed return of the hemodynamics to normal following pericardiectomy when ventricular volumes, wall

stress and stroke work are returned to normal. Dines and colleagues³⁰ selected a specific group of patients whose constriction was so severe that they died and were studied at autopsy.



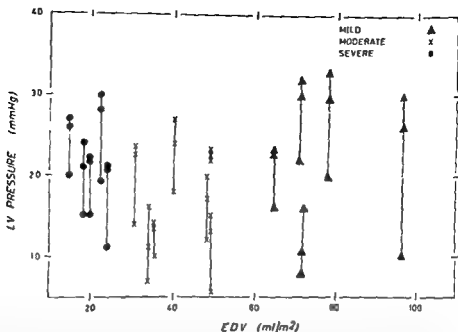


Fig 13 Relationship between LV end diastolic volume (EDV) and the diastolic LV pressures. The O point H point and EDP are shown for each patient and the values in each patient are joined by a vertical line. The EDP was similar in all the patients but patients with mild disease had wide diastolic pulse pressures (ΔP). This was small in patients with severe constriction and a small EDV.

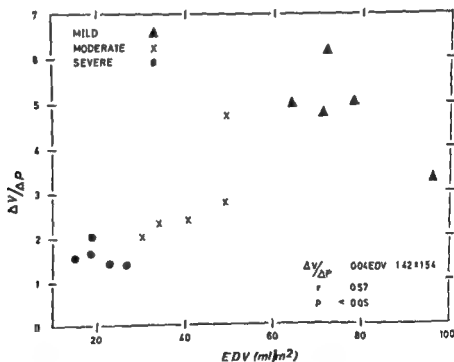


Fig 14 Simple linear relationship between LV distensibility index ($\Delta V/\Delta P$) and LV end diastolic volume (EDV). Patients with severe disease had a small EDV and a low distensibility index.

length, but fractional shortening is normal in patients in whom the ventricle is extrinsically splinted so that the crumpled compressed fibers operate at a mechanical disadvantage. The fibers cannot be stretched to a normal end diastolic length and operate on the lower part of the ascend-

ing limb of the Frank-Starling curve. The high EDP is a consequence of extrinsic compression and the fibers are underloaded despite the high filling pressure. Theoretically the fibers need to generate less tension and perform less work, but the constricting shell which infiltrates the epicardium also

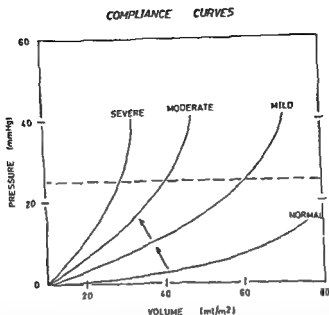


Fig. 17 Diastolic pressure volume curves for normal subjects and for patients with mild moderate and severe restriction of the ventriculo-pericardial system. At a critical venous filling pressure (dashed line) important clinical symptoms occur and the patients seek medical care regardless of their end-diastolic volume and compliance.

in the LV but this is not related to the EDV. Every diastolic pressure volume curve is exponential and the rate of pressure rise during ventricular filling depends on four factors: the slope of the curve, the level of the diastolic pressure, the end diastolic volume, and the absolute volume change. At high diastolic pressures, the curves have a steep slope so that changes in pressure and stroke volume overshadow the effect of changes in compliance or end diastolic volume.

There may also be time dependent changes due to viscous inertial or plastic properties of the pericardium. We have not measured these changes in the present study. Similarly, we have no information on the influence of the high end-diastolic pressure on coronary blood flow, but we presume it to be normal in the absence of coronary artery disease.

Summary

Left ventricular function was studied in systole and diastole in 30 patients with constrictive pericarditis. Left ventricular end-diastolic volume was used to divide the patients into three arbitrary groups: severe constriction ($EDV < 25 \text{ ml/M}^2$),

moderate constriction ($EDV 25 \text{ to } 50 \text{ ml/M}^2$), and mild constriction ($EDV > 50 \text{ ml/M}^2$).

The patients had high ventricular diastolic and venous filling pressures (mean $LVEDP = 23 \pm 7 \text{ mm Hg}$, mean $RVEDP = 20 \pm 7 \text{ mm Hg}$). Measurements related to absolute fiber shortening (stroke index, stroke work index, and left ventricular ejection rate) were reduced and linearly related to the degree of constriction as assessed by the end-diastolic volume.

Measurements of relative fiber shortening or lengthening (ejection and filling fraction and circumferential fiber shortening) were normal despite great reduction in ventricular volumes.

Velocity measurements (peak LV dp/dt and mean velocity of circumferential fiber shortening) were normal or slightly reduced.

These changes were reflected in the systolic time interval measurements: pre-ejection phase, left ventricular ejection time, and the ratio $PEP/LVET$.

Diastolic function of the ventricle was abnormal: the distensibility index of the ventriculo-pericardial system ($\Delta V/\Delta P$) was low and the passive elastic modulus increased. The change in compliance corre-

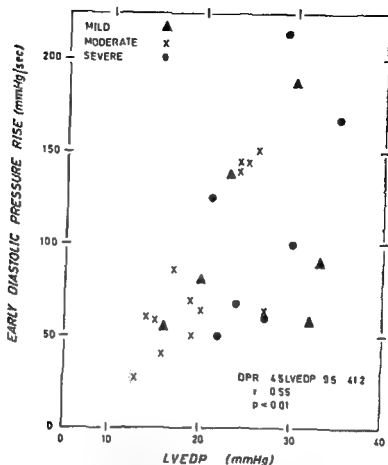


Fig 16 Relationship between the rate of pressure rise in early diastole (DPR) and LV end diastolic pressure (EDP). DPR was related to the LVEDP regardless of the severity of constriction assessed by angiography.

$$\Delta P/\Delta V = a\bar{P} + b$$

The measurement does not need measured angiographic volumes and expresses compliance in terms of changes in intraventricular diastolic pressure (ΔP) and stroke volume (ΔV). The passive elastic modulus was abnormal in constrictive pericarditis and increased with the severity of constriction. It is noteworthy that in severe constriction $\Delta P/\Delta V$ and the passive elastic modulus of the pericardium were lower than corresponding measurements of diastolic function of the myocardium in a group of patients with acute myocardial infarction.²¹

Our concept of the diastolic function of the ventricle in health and disease is shown schematically in Fig 17. There appears to be a family of diastolic pressure-volume curves for normal subjects and for patients with cardiac disease. Every patient is in a condition of circulatory equilibrium which is determined at any instant by the degree of constriction, the ventriculo-pericardial compliance, the stroke

output and the venous filling pressure. As the disease progresses and compliance falls, the patient moves from curve to curve. Important signs and symptoms of systemic venous hypertension become clinically apparent at a critical venous pressure and the patient seeks medical care regardless of the severity of constriction, the abnormality of compliance and the reduction in ventricular volume. This accounts for the similarity in the ventricular diastolic pressures in this group of patients. We did not investigate patients with mild constrictive pericarditis and it is possible that in the early phase of the disease the diastolic filling pressures which are much lower may be related to the severity of constriction. In established disease the diastolic pressures are similar; they do not correlate with the degree of constriction as assessed by angiocardiography and cannot be used to differentiate the severity of the constricting process.

Patients with a high EDP also have a rapid rate of pressure rise in early diastole.

- 33 Chatterjee K, Sacco M, Sutton, G C and Miller G A H. Assessment of left ventricular function by single plane cineangiographic volume analysis. *Br Heart J* 33:565 1971
- 34 Spotnitz H M, Sonnenblick E H and Spiro D. Relation of ultrastructure to function in the intact heart: sarcomere structure relative to pressure-volume curves of intact left ventricles of dog and cat. *Circ. Res* 18:49 1966
- 35 Noble M I M, Milne E N C, Goerke R J, Carlson E, Domenech R J, Saunders K H and Hoffman J I E. Left ventricular filling and diastolic pressure-volume relations in the conscious dog. *Circ Res* 24:269 1969
- 36 Monroe R G, Gamble W J, LaFarge C G, Kumar A E and Manasek F J. Left ventricular performance at high end-diastolic pressures in isolated perfused dog hearts. *Circ Res* 26: 1970
- 37 Fowler N B. *Cardiac diagnosis*. New York 1968. Harper & Row Publishers
- 38 Personal data
- 39 Barratt Boyes B G and Wood E H. Cardiac output and related measurements and pressure values in the right heart and associated vessels together with an analysis of the haemodynamic response to the inhalation of high oxygen mixtures in healthy subjects. *J Lab Clin Med* 51:72 1958
- 40 Kennedy J W, Baxley W A, Figley M M, Dodge H T and Blackmon J E. Quantitative angiocardiology I. The normal left ventricle in man. *Circulation* 34:272 1966
- 41 Snell R E and Luchsinger P C. Determination of the external work and power of the left ventricle in intact man. *AM HEART J* 69:529 1965
- 42 Levine H J, Neill W A, Wagman H J, Krasnow N and Gorlin R. The effect of exercise on mean left ventricular ejection rate in man. *J Clin Invest* 41:1050 1962

lated with the degree of constriction and there was a linear relationship between compliance and EDV

The ventricle was unloaded despite the high filling pressure and stroke work index was reduced extrinsic compression raised the diastolic pressure and reduced left ventricular volumes

REFERENCES

- Harvey R M Ferrer I, Cathcart R T, Richards D W and Courmand A Mechanical and myocardial factors in chronic constrictive pericarditis *Circulation* 8:695 1953
- Harrison E C Crawford D W and Lau F Y K Sequential left ventricular function studies before and after pericardiectomy for constrictive pericarditis *Am J Cardiol* 26:319 1970
- Vogel J H K Horgan J A and Strahl C L Left ventricular dysfunction in chronic constrictive pericarditis *Chest* 58:484 1971
- Dalton J C Pearson R J Jr and White P D Constrictive pericarditis: a review and long term follow up of 78 cases *Ann Intern Med* 45:445 1956
- Deterling R A Jr and Humphreys G H Factors in the etiology of constrictive pericarditis *Circulation* 12:130 1955
- Schrire V Experience with pericarditis at Groote Schuur Hospital Cape Town: an analysis of one hundred and sixty cases over a 6 year period *S Afr Med J* 33:810 1959
- Wood P Chronic constrictive pericarditis *Am J Cardiol* 7:148 1961
- Scheepers G W H Tuberculous pericarditis *Am J Cardiol* 9:148 1962
- Lewis B S van der Horst R L and Gotsman M S Diagnostic electrocardiographic patterns in Bantu myocardial pathology and constrictive pericarditis *S Afr Med J* 45:1110 1971
- Cortes F M The pericardium and its disorders ed 1 Springfield Ill 1971 Charles C Thomas Publisher
- Gotsman M S le Roux B T Rogers N M A van der Horst R L and Winship W S Immediate haemodynamic results of pericardiectomy *S Afr Med J* 46:1 1972
- Weissler A M Peeler R G and Roehll W H Relationship between left ventricular ejection time stroke volume and heart rate in normal individuals and patients with cardiovascular disease *Am Heart J* 62:367 1961
- Weissler A M Harris W S and Schoenfeld C D Systolic time intervals in heart failure in man *Circulation* 37:149 1968
- Weissler A M and Garrard C L Jr Systolic time intervals in cardiac disease (I) *Mod Concepts Cardiovasc Dis* 40:1 1971
- Weissler A M and Garrard C L Jr Systolic time intervals in cardiac disease II *Mod Concepts Cardiovasc Dis* 40:5 1971
- Garrard C L Jr Weissler A M and Dodge H T Relationships of alterations in systolic time intervals to ejection fraction in patients with cardiac disease *Circulation* 42:455 1970
- Weissler A M and Schoenfeld C D Effect of digitalis on systolic time intervals in heart failure *Am J Med Sci* 259:4 1970
- Greene D G Carlisle R Grant C and Bunnell I L Estimation of left ventricular volume by one-plane cineangiography *Circulation* 33:61 1967
- Carleton R A Change in left ventricular volume during angiocardiology *Am J Cardiol* 27:460 1971
- Gaasch W H Battle W E Oboler A A Banas J S and Levine H J Left ventricular stress and compliance in man *Circulation* 45:746 1972
- Diamond G and Forrester J S Effect of coronary artery disease and acute myocardial infarction on left ventricular compliance in man *Circulation* 45:11 1972
- Diamond G Forrester J B Hargis J Parmley W W Danzig R and Swan H J C Diastolic pressure-volume relationship in the canine left ventricle *Circ Res* 29:167 1971
- Sawyer C G Burwell C S Dexter L Eppinger E C Goodale W T Gorlin R Harken D E and Haynes F W Chronic constrictive pericarditis further consideration of the pathologic physiology of the disease *Am Heart J* 44:207 1952
- Wilson R H Hoseth W Sadoff C and Dempsey M E Pathologic physiology and diagnostic significance of the pressure pulse tracings in the heart in patients with constrictive pericarditis and pericardial effusion *Am Heart J* 48:671 1954
- Kesteloot H and Denef B Value of reference tracings in diagnosis and assessment of constrictive epicarditis and pericarditis *Br Heart J* 32:675 1970
- Shabetai R Fowler N O and Guntheroth W G The hemodynamics of cardiac tamponade and constrictive pericarditis *Am J Cardiol* 26:480 1970
- Hancock E N Subacute effusive-constrictive pericarditis *Circulation* 43:183 1971
- Kloster F E Crislip R L Bristow J D Herr R H Ritzmann L W and Griswold H E Hemodynamic studies following pericardiectomy for constrictive pericarditis *Circulation* 32:415 1965
- Albers W H Hugenholtz P G and Nadas A S Constrictive pericarditis and atrial septal defect secundum type *Am J Cardiol* 23:850 1969
- Dines D E Edwards J E and Burchell H H Myocardial atrophy in constrictive pericarditis *Mayo Clin Proc* 33:93 1958
- Roberts J T and Beck C H The effect of chronic cardiac compression on the size of heart muscle fibers *Am Heart J* 22:314 1941
- Chatterjee K Saccor M Sutton G C and Miller G A H Angiographic assessment of left ventricular function in patients with ischaemic heart disease without clinical heart failure *Br Heart J* 33:559 1971

used in this investigation are presented elsewhere¹¹⁾

Methods

The corrected orthogonal leads of 120 normal subjects were recorded on magnetic tape using the axial lead system of McFee and Parungao. In order to determine if these subjects were free of heart disease each was given a clinical examination (Table I) including measurement of his blood pressure cardiac auscultation and evaluation by palpation and percussion. Histories of these subjects were also studied in an effort to detect any previous cardiovascular problems.

The details of procedures utilized in recording and signal processing were reported in an earlier paper.¹² A total of 63 measurements were made from the digitized wave forms. The results of the analysis of these measurements are reported in Tables II through V.

Thirty six of the 63 distributions of the measurements differed significantly ($p < 0.01$) from normality some because of skewness alone and others because of skewness and kurtosis. Logarithmic and linear transformations were performed on those measurements whose distributions were skewed. The means and confidence limits of the transformed data were then computed. These estimates were then retransformed to the linear scale and the results were reported. Twenty five of the 36 nonnormal distributions thus transformed were not significantly different ($p < 0.01$) from the Gaussian curve.

Two of the remaining 11 non Gaussian distributions were significantly different ($p < 0.01$) from normality because of kurtosis. The direction of terminal deflection of the QRS loop in the frontal plane was platykurtic i.e. widely scattered and somewhat flat the mean is reported in Table III. The distribution of the direction of the 40 msec vector of the QRS loop in

Table I Distribution of subjects by age and sex

| Age (yrs) | Male | Female | Total |
|-----------|------|--------|-------|
| 15 to 24 | 22 | 25 | 47 |
| 25 to 34 | 34 | 18 | 52 |
| 35 to 44 | 3 | 9 | 12 |
| 45 to 54 | 2 | 6 | 8 |
| 55 to 64 | 0 | 1 | 1 |
| Total | 61 | 59 | 120 |

the left sagittal plane was found to be leptokurtic (highly peaked). The mean and 95 per cent confidence limits of this distribution are reported however since it was felt that this information was still valuable and consistent with the data.

Transformation of the 9 remaining non Gaussian distributions (the direction of the maximum P vector in all three planes the direction of terminal deflection of the QRS loop in the horizontal plane the magnitude of the 60 msec vector in the QRS loop QRS duration the magnitude of the ST vector in the left sagittal plane and the direction of the maximum T_r vector in both the frontal and left sagittal planes) did not eliminate the problems of non normality therefore the medians and ranges of these data are reported in Table II.

The recommendations for reference axes of the Committee on Electrocardiography of the American Heart Association were followed with one exception. The zero position on the left sagittal reference frame was transposed to the patient's anterior. The numerical data thus obtained are the same as if the right sagittal reference frame had been used and can be compared readily to the normal values reported by Chou and Helm¹³ and Macfarlane and associates.¹⁴

Results

The statistics which summarize the data are shown in Tables II through V. An inspection of these tables will reveal that some of the distributions were Gaussian in form and required no transformation however a majority of the measurements were non Gaussian. The logarithmic transformation procedure made most of these non

If the d t were ll wed t th right b t sy t t
a d d d t were that ll l be g t than se o
d the log r h m were then t ken. If th dat were
k wed t th left th d ta were d t m l t p l d by -1
and the tra of m d t th name m ne Retra of rma
tio of the p rmetre stin te would then l d t takt g
the t log f th m ud limits, bt cti g sy
t t th t h d bee d d d d d m hply g by -1
if the d ta were kewed to the left

The McFee-Parungao (axial) vectorcardiogram in normal subjects

G Daniel Copeland, M D

Ann B McEachran B S

Herbert W Smith, Ph D

Daniel A Brody, M D

Memphis, Tenn

The McFee Parungao (axial) vectorcardiographic lead system¹ has been shown to have theoretical advantages over other vectorcardiographic lead systems now in use.²⁻⁴ Recently, Lawrie and Macfarlane⁵ reported that in clinical application, the axial lead system correlated more favorably with an ideal orthogonal lead system than did the widely used Frank system. Macfarlane⁶ modified the axial lead system by means of the Brody Arzbaecher lead strength equalization and published the results of extensive clinical trials using this modification.⁷⁻⁹ Gabor and White¹⁰ and Ainger¹¹⁻¹³ studied several lead systems in infants including the axial, and the results of these investigations continue to appear in the literature. The purpose of this paper is to present estimates of the parameters of the unmodified axial vectorcardiograms in normal adults.

Comparisons among lead systems are difficult for two reasons. First because many electrocardiographic measurements are non Gaussian in distribution, the determination of means and 95 per cent con-

fidence limits for given measurements in electrocardiography and vectorcardiography has been difficult. To avoid this problem some authors present ranges while others disregard the distributions of the data and calculate 95 per cent confidence limits. Second, comparison of parameter estimates is sometimes difficult because data are measured in different ways. For example some workers¹³⁻¹⁶ report values measured at the change of direction of the vector loop while most workers in this country and Europe report values obtained at various timed intervals after the onset of the loop. Downs and Liebman,¹⁷ and Ainger¹¹⁻¹³ indicate that linear methods of reporting and analyzing angular data are incorrect and treat vector quantities as points on a circle or a sphere.

In this study, yet another method for presentation of the vectorcardiographic data is used: logarithmic transformations have been applied to those planar vectorcardiographic values whose distributions are not Gaussian. (Descriptive statistics for some of the untransformed data

From the Division of Clinical Physiology, University of Tennessee Medical Units, Memphis, Tenn.
Supported by Grants HL-01362, HL-14032, and HL-09495 of the National Institutes of Health, United States Public Health Service.

Received for publication Aug. 30, 1972.

Reprint requests to Daniel A. Brody, M.D., University of Tennessee Medical Units, Division of Clinical Physiology, 951 Court Ave., Memphis, Tenn. 38103.

Table IV The direction and magnitude of the junctional (S T) vector in normal subjects

| S T vector | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|---------------------|------------------|-----------------|---------------|-----------------|---------------------|-----------------|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Magnitude (mv) | 0.50 | 0.24 to 1.01 | 0.50 | 0.27 to 0.93† | 0.38 | 0.07 to 0.91† |
| Direction (degrees) | +29 | -4 to +61 | +31 | +10 to 52 | +49 | +5 to +93 |

Me and 95 per cent confidence limits computed using logarithmic transformation
 †Median and range reported

Table V The direction and magnitude of the maximum T vector in normal subjects

| T vector | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|-------------------------------|------------------|-----------------|---------------|-----------------|---------------------|-----------------|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Maximum T vector | | | | | | |
| Magnitude (mv) | 0.53 | 0.25 to 1.15 | 0.51 | 0.25 to 1.02 | 0.39 | 0.14 to 1.10 |
| Direction (degrees) | +33 | 0 to +66 | +31 | +9 to +53 | +42 | +2 to +83 |
| T duration (msec) | 284 | 232 to 336 | | | | |
| Maximum T _P vector | | | | | | |
| Magnitude (mv) | 0.03 | 0.01 to 0.07 | 0.04 | 0.02 to 0.09 | 0.04 | 0.01 to 0.09 |
| Direction (degrees) | -119 | +133 to -11 | -110† | +47 to 0† | -118† | -177 to +62† |

Me and 95 per cent confidence limits computed by using logarithmic transformation
 †Median and range reported

Table VI Lead strengths of the modified and original axial lead systems also the Frank lead system

| System | X | Y | Z |
|----------------|------|------|------------|
| Axial | 1.61 | 1.48 | 2.24 mv/cm |
| Modified axial | 1.48 | 1.48 | 1.48 |
| Frank | 1.45 | 1.40 | 1.37 |

in comparison with the QRS and T loops. Generally the P loop tends to be inscribed counterclockwise in the horizontal and left sagittal planes. In the frontal plane the loop is even more variable and a figure of eight configuration is frequently seen.

The QRS loop. Nearly all the QRS loops were inscribed counterclockwise in the horizontal and left sagittal planes (95 per cent in each plane). In these two planes

those loops (7 per cent in each plane) which present a figure-of-eight morphology were predominantly counterclockwise. The initial portion of those loops was always counterclockwise in this series and the change of direction which produced the figure-of-eight occurred distal from the origin of the loop and near the site of the maximum instantaneous vector. Details of these loops are presented in Table III. Generally the values are larger than similar values reported with the Frank system or with the modified axial lead system; this is especially true of the 40 millisecond to 60 millisecond and maximum instantaneous vectors.

The direction of the initial deflection. of interest in the three planes. In the horizontal plane the mean direction was 106 degrees (anterior and slightly to the right) with limits from 41 to 158 degrees. It can thus be said that the initial vector in the horizontal plane always moves in an an-

Table II The direction and magnitude of the maximum P vector in normal subjects

| Parameter | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|---------------------------|------------------|-----------------|---------------|-----------------|---------------------|-----------------|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Maximum P vector | | | | | | |
| Magnitude (mv) | 0.08 | 0.05 to 0.14 | 0.13 | 0.06 to 0.2 | 0.12 | 0.04 to 0.21 |
| Time of occurrence (msec) | 64 | 35 to 94 | 55 | 32 to 78 | 50 | 23 to 78 |
| Direction (degrees) | -5† | -100 to +161† | +66† | -46 to +107† | +82† | -165 to +119† |
| P duration (msec) | 94.0* | 75 to 118* | | | | |

*Mean and 95 per cent confidence limits computed using logarithmic transformation

†Median and range reported

Table III Characteristics of the normal QRS loop

| Characteristic | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|---|------------------|--|---------------|--|---------------------|--|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Maximum QRS vector | | | | | | |
| Magnitude (mv) | 1.61* | 0.94 to 2.78* | 1.97 | 1.01 to 2.92 | 1.48 | 0.62 to 2.34 |
| Direction (degrees) | -17* | -120 to +38* | +39 | +10 to +69 | +119 | +40 to -161 |
| Time of occurrence of max QRS vector (msec) | 46* | 34 to 63* | 43* | 35 to 54* | 47* | 34 to 64* |
| 20 msec vector | | | | | | |
| Magnitude (mv) | 0.43* | 0.16 to 1.13* | 0.16* | 0.03 to 0.76* | 0.40* | 0.15 to 1.08* |
| Direction (degrees) | +78 | +30 to +126 | 0 | -163 to +163 | +4 | -37 to +45 |
| 40 msec vector | | | | | | |
| Magnitude (mv) | 1.42 | 0.52 to 2.32 | 1.68 | 0.64 to 2.73 | 1.16 | 0.33 to 1.98 |
| Direction (degrees) | -5* | -73 to +38 | +38 | +5 to +71 | +98* | +23 to +156* |
| 60 msec vector | | | | | | |
| Magnitude (mv) | 0.79† | 0.13 to 2.14† | 0.31* | 0.06 to 1.61* | 0.72* | 0.23 to 2.28* |
| Direction (degrees) | -89 | -137 to -26* | -7 | All quadrants | +178* | +114 to -129 |
| Direction of initial deflection (degrees) | +106* | +41 to +158 | -138* | +81 to -31 | +20* | -69 to +56* |
| Direction of terminal deflection (degrees) | -95† | All quadrants | -43* | All quadrants-169 | | +107 to -85 |
| Direction of inscription | | 93% Counterclockwise 7% Figure of eight | | 36% Clockwise 43% Figure of eight 21% Counterclockwise | | 7% Figure of eight 93% Counterclockwise |
| QRS duration (msec) | 94† | 70 to 142† | | | | |

*Mean and 95 per cent confidence limits computed using logarithmic transformation

†Median and range reported

normal distributions Gaussian (these are indicated by one asterisk in the tables) and permitted the computation of valid 95 per cent confidence limits. A third group of data could not be transformed to a normal distribution (indicated by a dagger). For this group only ranges and medians are reported.

The P loop The atrial loop does not close because the atrial T deflection, which is normally opposite in sign from the P wave in orthogonal and scalar leads begins before the end of the atrial depolarization process. The P loop is very irregular in contour and is so small that considerable amplification is required to visualize it well.

Table IV The direction and magnitude of the junctional (ST) vector in normal subjects

| S-T vector | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|---------------------|------------------|-----------------|---------------|-----------------|---------------------|-----------------|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Magnitude (mv) | 0.50 | 0.24 to 1.01 | 0.50 | 0.27 to 0.93† | 0.38 | 0.07 to 0.91† |
| Direction (degrees) | +29 | -4 to +61 | +31 | +10 to 52 | +49 | +5 to +93 |

Mean and 95 per cent confidence limits computed using logarithmic transformation.

†Median and range reported

Table V The direction and magnitude of the maximum T vector in normal subjects

| T vector | Horizontal plane | | Frontal plane | | Left sagittal plane | |
|-------------------------------|------------------|-----------------|---------------|-----------------|---------------------|-----------------|
| | Mean | 95% Conf limits | Mean | 95% Conf limits | Mean | 95% Conf limits |
| Maximum T vector | | | | | | |
| Magnitude (mv) | 0.53 | 0.25 to 1.15 | 0.51 | 0.25 to 1.02 | 0.39 | 0.14 to 1.10 |
| Direction (degrees) | +33 | 0 to +66 | +31 | +9 to +53 | +42 | +2 to +83 |
| T duration (msec) | 284 | 232 to 336 | | | | |
| Maximum T ₂ vector | | | | | | |
| Magnitude (mv) | 0.03 | 0.01 to 0.07 | 0.04 | 0.01 to 0.09 | 0.04 | 0.02 to 0.09 |
| Direction (degrees) | -119 | +133 to -11 | -110† | +47 to 0† | -118† | -177 to +62† |

Mean and 95 per cent confidence limits computed by using the logarithmic transformation.

†Median and range reported

Table VI Lead strengths of the modified and original axial lead systems also the Frank lead system

| System | X | Y | Z |
|----------------|------|------|------------|
| Axial | 1.61 | 1.48 | 2.24 mv/cm |
| Modified axial | 1.48 | 1.48 | 1.48 |
| Frank | 1.45 | 1.40 | 1.37 |

in comparison with the QRS and T loops. Generally the P loop tends to be inscribed counterclockwise in the horizontal and left sagittal planes. In the frontal plane the loop is even more variable and a figure of eight configuration is frequently seen.

The QRS loop. Nearly all the QRS loops were inscribed counterclockwise in the horizontal and left sagittal planes (90 per cent in each plane). In these two planes

those loops (7 per cent in each plane) which present a figure-of-eight morphology were predominantly counterclockwise. The initial portion of those loops was always counterclockwise in this series and the change of direction which produced the figure-of-eight occurred distal from the origin of the loop and near the site of the maximum instantaneous vector. Details of these loops are presented in Table III. Generally the values are larger than similar values reported with the Frank system or with the modified axial lead system; this is especially true of the 40 millisecond, 60 millisecond and maximum instantaneous vectors.

The direction of the initial deflection is of interest in the three planes. In the horizontal plane the mean direction was 106 degrees (anterior and slightly to the right) with limits from 41 to 158 degrees. It can thus be said that the initial vector in the horizontal plane always moves in an an-

Table VII *P loop, maximum P vector*

| Source | Transverse plane | | |
|-----------------------------------|---------------------|----------------|-------------|
| | Direction (degrees) | Magnitude (mv) | Time (msec) |
| Chou and Helm (Frank) | | | |
| Mean | -5 | 0.07 | |
| 95% Ranges | -50 to 60 | 0.04 to 0.10 | |
| Macfarlane et al (Modified axial) | | | |
| (a) Up through 14 yrs | | | |
| Mean and S D | 21 ± 50 | 0.09 ± 0.03 | |
| 96% Ranges | -46 to 101 | 0.06 to 0.14 | |
| (b) 15 to 29 yrs | | | |
| Mean and S D | 25 ± 53 | 0.08 ± 0.03 | |
| 96% Ranges | 2 to 165 | 0.03 to 0.16 | |
| (c) > 29 yrs of age | | | |
| Mean and S D | 3 ± 54 | 0.08 ± 0.03 | |
| 96% Ranges | 2 to 103 | 0.04 to 0.14 | |
| Present study (Axial) | | | |
| Mean | -5† | 0.08 | 64 |
| 95% Conf limits | -100 to 161† | 0.05 to 0.14 | 35 to 94 |
| P duration (msec) | 94.0* | 75 to 118* | |

*Mean and 95 per cent confidence limits were computed using logarithmic transformation

†Median and range reported

Table VIII *QRS loop, maximum QRS vector*

| Source | Transverse plane | | |
|------------------------------------|---------------------|----------------|-------------|
| | Direction (degrees) | Magnitude (mv) | Time (msec) |
| Chou and Helm (Frank) | | | |
| Mean | -10 | 1.3 | 38 |
| 96% Ranges | -80 to 20 | 0.85 to 1.95 | 30 to 48 |
| Macfarlane et al (Corrected axial) | | | |
| (a) Up through 14 yrs | | | |
| Mean and S D | -25 ± 56 | 1.34 ± 0.45 | |
| 96% Ranges | -125 to 40 | 0.81 to 2.07 | |
| (b) 15 to 29 yrs | | | |
| Mean and S D | -11 ± 26 | 1.15 ± 0.34 | |
| 96% Ranges | -88 to 30 | 0.52 to 1.98 | |
| (c) 30 yrs and older | | | |
| Mean and S D | -5 ± 24 | 1.12 ± 0.33 | |
| 96% Ranges | -82 to 23 | 0.59 to 1.89 | |
| Present study (Axial) | | | |
| Mean | -17* | 1.61* | 46* |
| 95% Conf limits | -120 to 38* | 0.94 to 2.78* | 34 to 63* |

*Mean and 95 per cent confidence limits computed by use of logarithmic transformation

terior direction but may be either to the left or right. In the frontal plane the mean direction of the initial deflection was -138 degrees, with limits of -31 to 81 degrees confirming the left to right direction tend

ency and the greater variability of measurements in the frontal plane. In the left sagittal plane the mean direction of the initial vector was 20 degrees with limits from -69 to 56 degrees that initial

| Sagittal plane | | | Frontal plane | | |
|-----------------------|-----------------------------|----------------|----------------------|-----------------------------|----------------|
| Direction (degrees) | Magnitude (mv) | Time (msec) | Direction (degrees) | Magnitude (mv) | Time (msec) |
| 85 50 to 110 | 0.12 0.04 to 0.18 | | 65 15 to 90 | 0.12 0.06 to 0.20 | |
| 47 ± 64 -60 to 108 | 0.12 ± 0.04 0.04 to 0.19 | | 40 ± 45 -20 to 70 | 0.13 ± 0.04 0.09 to 0.17 | |
| 64 ± 43 41 to 108 | 0.13 ± 0.06 0.04 to 0.25 | | 49 ± 42 27 to 99 | 0.13 ± 0.06 0.05 to 0.25 | |
| 86 ± 41 -129 to 1 | 0.14 ± 0.06 0.06 to 0.27 | | 62 ± 38 -8 to 99 | 0.15 ± 0.05 0.06 to 0.24 | |
| 82† -165 to 119† | 0.12 0.04 to 0.21 | 50 23 to 78 | 66† -46 to 107† | 0.13 0.06 to 0.2 | 55 32 to 78 |

| Sagittal plane | | | Frontal plane | | |
|------------------------|-----------------------------|----------------|---------------------|-----------------------------|-----------------|
| Direction (degrees) | Magnitude (mv) | Time (msec) | Direction (degrees) | Magnitude (mv) | Time (msec) |
| 100 50 to 165 | 1.0 0.3 to 1.9 | Same Same | 35 10 to 65 | 1.5 0.9 to 2.2 | Same Same |
| 102 ± 30 52 to 155 | 1.33 ± 0.40 0.70 to 1.98 | | 43 ± 14 27 to 62 | 1.74 ± 0.54 0.99 to 2.41 | |
| 110 ± 33 64 to -151 | 1.24 ± 0.45 0.59 to 2.16 | | 45 ± 14 17 to 74 | 1.61 ± 0.44 1.00 to 2.21 | |
| 109 ± 42 49 to -50 | 0.91 ± 0.32 0.47 to 1.63 | | 35 ± 19 0 to 63 | 1.37 ± 0.38 0.79 to 2.29 | |
| 119 -161 to 40 | 1.48 0.62 to 2.34 | 47 34 to 64 | 39 10 to 69 | 1.97 1.01 to 2.92 | 43* 35 to 54 |

deflection should be anterior and may be directed either up or down.

The ST (junctional) vector and the T vector. The statistics for these two vectors are included in Tables IV and V to show

their relationship to the vectorcardiographic loop as well as to the scalar electrocardiogram. The T loop is slowly inscribed. It is generally smooth in contour as is the QRS loop. The T loop is nearly always in

Table IX Normal QRS loop

| Source | Transverse plane | | Sagittal plane | | Frontal plane | |
|-----------------------|---------------------|----------------|---------------------|----------------|---------------------|----------------|
| | Direction (degrees) | Magnitude (mV) | Direction (degrees) | Magnitude (mV) | Direction (degrees) | Magnitude (mV) |
| 20 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | 55 | 0.40 | 15 | 0.30 | 20 | 0.30 |
| 95% Ranges | 0 to 120 | 0.15 to 0.75 | -30 to 75 | 0.1 to 0.55 | All quadrants | 0.05 to 0.7 |
| Present study (Axial) | | | | | | |
| Mean | 78 | 0.43* | 4 | 0.40* | 0 | 0.16* |
| 95% Conf limits | 30 to 126 | 0.16 to 1.13* | -37 to 45 | 0.15 to 1.08* | -163 to 163 | 0.03 to 0.76* |
| 40 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | -20 | 1.15 | 110 | 0.9 | 35 | 1.25 |
| 95% Ranges | -90 to 25 | 0.55 to 1.9 | 60 to 170 | 0.25 to 1.8 | -10 to 70 | 0.4 to 2.2 |
| Present study (Axial) | | | | | | |
| Mean | -5* | 1.42 | 98* | 1.16 | 38 | 1.88 |
| 95% Conf limits | -73 to 38* | 0.52 to 2.32 | 23 to 156* | 0.33 to 1.98 | 5 to 71 | 0.64 to 2.73 |
| 60 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | -90 | 0.43 | 160 | 0.55 | 85 | 0.25 |
| 95% Ranges | -125 to -35 | 0.0 to 0.9 | 115 to -40 | 0.1 to 1.0 | All quadrants | 0.0 to 0.6 |
| Present study (Axial) | | | | | | |
| Mean | -89* | 0.79† | 178* | 0.72* | -7 | 0.31* |
| 95% Conf limits | -137 to -26* | 0.13 to 2.14† | 114 to -129* | 0.23 to 2.28* | All quadrants | 0.06 to 1.61* |

*Mean and 95 per cent confidence limits computed using logarithmic transformation

†Median and range reported

scribed in the same direction as the QRS loop in the normal individual however there was an occasional subject in the sample for whom this was not the case

Discussion

Presented in Table VI are the lead strengths in millivolts per centimeter for the axial modified axial and Frank lead systems. In Tables VII through X the data obtained in the present study are compared with the data reported by Chou and Helm¹⁹ (Frank lead system) and by Macfarlane and co-workers⁷ (modified axial lead system). Although the present study uses the left sagittal reference frame while the other two studies use the right sagittal the numerical data should be comparable since the polarity of the abscissa is reversed. The data in this report and in Macfarlane's were translated to allow degrees to be re-

ported to plus 180 degrees and minus 180 degrees.

The data in Tables VII, VIII, and X reflect Macfarlane's three groups of data: (a) up through 14 years, (b) 15 through 29 years, and (c) 30 years and over.

Chou and Helm reported their data with means and 95 per cent ranges while Macfarlane and colleagues reported means, standard deviations, and 96 per cent ranges.

It is apparent that the greater lead strengths of the axial (McFee Parungao) lead system are more manifest in the QRS loop than in either the P or T loop. This is in accordance with the observed greater size or more rotund appearance of the axial QRS loops. The observation that the 40 millisecond, 60 millisecond, and maximum instantaneous vectors are all substantially greater in the axial system is

Table X *T loop maximum T vector*

| Source | Transverse plane | | Sagittal plane | | Frontal plane | |
|------------------------------------|---------------------|----------------|---------------------|----------------|---------------------|----------------|
| | Direction (degrees) | Magnitude (mv) | Direction (degrees) | Magnitude (mv) | Direction (degrees) | Magnitude (mv) |
| Chou and Helm (Frank) | | | | | | |
| Mean | 35 | 0.5 | 45 | 0.4 | 35 | 0.5 |
| 95% Ranges | 0 to 65 | 0.25 to 0.75 | 20 to 90 | 0.2 to 0.7 | 20 to 55 | 0.25 to 0.75 |
| Macfarlane et al (Corrected axial) | | | | | | |
| (a) Up through 14 yrs | | | | | | |
| Mean and S D | 12.5 ± 17 | 0.44 ± 0.21 | 76 ± 21 | 0.29 ± 0.17 | 32 ± 13 | 0.50 ± 0.23 |
| 96% Ranges | -5 to 36 | 0.24 to 0.83 | 46 to 106 | 0.14 to 0.59 | 15 to 54 | 0.26 to 0.94 |
| (b) 15 to 29 yrs | | | | | | |
| Mean and S D | 30 ± 21 | 0.35 ± 0.13 | 58 ± 23 | 0.32 ± 0.13 | 41 ± 20 | 0.41 ± 0.14 |
| 96% Ranges | -6 to 78 | 0.15 to 0.64 | 27 to 102 | 0.14 to 0.62 | 17 to 70 | 0.18 to 0.78 |
| (c) 30 yrs and older | | | | | | |
| Mean and S D | 29 ± 23 | 0.31 ± 0.11 | 59 ± 22 | 0.30 ± 0.11 | 42 ± 18 | 0.36 ± 0.12 |
| 96% Ranges | -3 to 71 | 0.13 to 0.56 | 25 to 92 | 0.14 to 0.54 | 15 to 68 | 0.16 to 0.59 |
| Present study (Axial) | | | | | | |
| Mean | 33 | 0.53 | 42 | 0.39 | 31 | 0.51 |
| 95% Conf limits | 0 to 66 | 0.25 to 1.15 | 2 to 83 | 0.14 to 1.10 | 9 to 53 | 0.25 to 1.02* |
| T duration (msec) | 284 | | | | | |
| | 232 to 336 | | | | | |

*Th 95 per cent confidence limits were computed by using the logarithmic transformation.

of obvious importance in deriving any voltage criteria for ventricular hypertrophy. The modified axial values fall generally between the axial and the Frank values and indeed the published modified axial loops closely resemble Frank loops rather than unmodified axial loops. As far as the planar projections of the maximum instantaneous P vector are concerned the mean magnitudes in all the reports are in close agreement (Table VII).

The 20, 40 and 60 millisecond vectors of the QRS in this study are compared in Table IX with those reported by Chou and Helm for the Frank system. With one exception the mean magnitude of the 20 millisecond vector in the frontal plane the mean values of the unmodified axial lead system were again larger than those of the Frank lead system.

Although transformation of nonnormal data is a fairly common statistical procedure²⁰ vectorcardiographers appear not to have used this method of obtaining means and 95 per cent confidence limits for skewed electrocardiographic distributions.

Failure to transform data whose distributions are skewed leads to incorrect confidence limits and the clinician could conclude from using these incorrect limits that a subject was outside normal limits when he was not.

Summary

Sixty-three parameter estimates from the McFee Parungao (axial) vectorcardiograms of 120 normal subjects were computed and tested for normality. Thirty-six of these distributions were found to differ significantly ($p < 0.01$) from Gaussian distribution. Logarithmic transformations were performed on those measurements whose distributions were nonnormal and it was found that all but nine distributions so treated did not differ significantly from the Gaussian distribution.

Thirty-nine of the 63 measurements were compared with electrocardiographic data from the modified axial and Frank lead systems. Lead strengths for the QRS vector in the axial system were larger than those for the other two systems. This difference

Table IX Normal QRS loop

| Source | Transverse plane | | Sagittal plane | | Frontal plane | |
|-----------------------|---------------------|----------------|---------------------|----------------|---------------------|----------------|
| | Direction (degrees) | Magnitude (mv) | Direction (degrees) | Magnitude (mv) | Direction (degrees) | Magnitude (mv) |
| 20 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | 55 | 0.40 | 15 | 0.30 | 20 | 0.30 |
| 95% Ranges | 0 to 120 | 0.15 to 0.75 | -30 to 75 | 0.1 to 0.55 | All quadrants | 0.05 to 0.7 |
| Present study (Axial) | | | | | | |
| Mean | 78 | 0.43* | 4 | 0.40* | 0 | 0.16* |
| 95% Conf lmts | 30 to 126 | 0.16 to 1.13* | -37 to 45 | 0.15 to 1.08* | -163 to 163 | 0.03 to 0.76* |
| 40 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | -20 | 1.15 | 110 | 0.9 | 35 | 1.25 |
| 95% Ranges | -90 to 25 | 0.55 to 1.9 | 60 to 170 | 0.25 to 1.8 | -10 to 70 | 0.4 to 2.2 |
| Present study (Axial) | | | | | | |
| Mean | -5° | 1.42 | 98° | 1.16 | 38 | 1.68 |
| 95% Conf lmts | -73 to 38* | 0.52 to 2.32 | 23 to 156* | 0.33 to 1.98 | 5 to 71 | 0.64 to 2.73 |
| 60 msec Vector | | | | | | |
| Chou and Helm (Frank) | | | | | | |
| Mean | -90 | 0.45 | 160 | 0.55 | 85 | 0.25 |
| 95% Ranges | -125 to -35 | 0.0 to 0.9 | 115 to -40 | 0.1 to 1.0 | All quadrants | 0.0 to 0.6 |
| Present study (Axial) | | | | | | |
| Mean | -89° | 0.79† | 178° | 0.72° | -7 | 0.31* |
| 95% Conf lmts | -137 to -26* | 0.13 to 2.14† | 114 to -129* | 0.23 to 2.28* | All quadrants | 0.06 to 1.61* |

*Mean and 95 per cent confidence limits computed using logarithmic transformation

†Median and range reported

scribed in the same direction as the QRS loop in the normal individual however there was an occasional subject in the sample for whom this was not the case

Discussion

Presented in Table VI are the lead strengths in millivolts per centimeter for the axial modified axial and Frank lead systems. In Tables VII through X the data obtained in the present study are compared with the data reported by Chou and Helm¹⁹ (Frank lead system) and by Macfarlane and co workers⁷ (modified axial lead system). Although the present study uses the left sagittal reference frame while the other two studies use the right sagittal, the numerical data should be comparable since the polarity of the abscissa is reversed. The data in this report and in Macfarlane's were translated to allow degrees to be re-

ported to plus 180 degrees and minus 180 degrees.

The data in Tables VII, VIII and X reflect Macfarlane's three groups of data (a) up through 14 years (b) 15 through 29 years and (c) 30 years and over.

Chou and Helm reported their data with means and 95 per cent ranges while Macfarlane and colleagues reported means, standard deviations and 96 per cent ranges.

It is apparent that the greater lead strengths of the axial (McFee Parungao) lead system are more manifest in the QRS loop than in either the P or T loop. This is in accordance with the observed greater size or more rotund appearance of the axial QRS loops. The observation that the 40 millisecond, 60 millisecond and maximum instantaneous vectors are all substantially greater in the axial system is

Experience with the coronary artery bypass graft in a community hospital

James N. Karnegis MD PhD
St Paul Minn

Although the coronary artery bypass graft (CABG) now accounts for about one half of all open heart operations¹ it remains an experimental procedure². Many of the reports which are available have come from a relatively limited number of large institutions where the technique has been developed and a large volume of work done. The future role of the operation will become better defined as a broader experience from more and different types of institutions is obtained. The purpose of this paper is to review the hospital course of patients undergoing CABG in a large private hospital.

Case material and results

Fifty five patients were submitted to the CABG procedure. Eleven died and 44 were discharged from the hospital giving an early mortality rate of 20 per cent. Table I compares some clinical features of the two groups. The average age of the survivors was 53 and of the nonsurvivors 63 years. Forty two of the 55 patients were men. The average clinical class using the New York Heart Association classification was 3.0 for those who lived and 3.5 for those who did not. Statistical analysis by the t test showed that there was no difference in

clinical class between the two groups at the 0.10 level of significance.

A history of congestive heart failure and probably also systemic hypertension occurred more frequently in the nonsurvivors. However the incidence of a history of diabetes mellitus, hyperlipidemia or a previous myocardial infarction was about the same in both groups.

Table II compares some angiographic features of the surviving and nonsurviving patients. All patients had at least one major coronary artery with a luminal narrowing of 50 per cent or more. The survivors had an average of 2.1 coronary arteries per patient demonstrating a luminal narrowing of at least 50 per cent compared with 2.4 for the nonsurvivors. The presence of mitral insufficiency and abnormalities of the wall of the left ventricle and of its contraction were frequent in both groups.

Table III shows that 77 grafts were done in the 44 survivors and 20 in the 11 nonsurvivors giving an average number of grafts per patient of 1.75 and 1.82 respectively. Statistical analysis by the t test showed that there was no difference in the average number of grafts per patient between the two groups at the 0.20 level of

From the Cardiac Diagnostic Clinic, The Charles T. Miller Hospital and the University of Minnesota Medical School, Minneapolis.
Received for publication Sept. 25, 1972.
Reprint requests to Dr. J. N. Karnegis, MD, The Charles T. Miller Hospital, 125 West College Avenue, St. Paul, MN 55102.

resulted in the axial systems having greater mean planar projections for the 40 millisecond 60 millisecond, and maximum instantaneous QRS vectors than either of the other two systems. There are few differences in P and T loops among the three, and the directions of inscription are similar for the three systems. As far as magnitude is concerned, the QRS values for the modified axial lead system fall between the values for axial and Frank lead systems.

REFERENCES

- McFee R and Parungao A. An orthogonal lead system for clinical electrocardiography. *AM HEART J* 61: 93 1961
- Brody D A and Arzbaecher R C. A comparative analysis of several corrected vectorcardiographic leads. *Circulation* 29: 533, 1964
- Brody D A and Arzbaecher R C. Intrinsic properties of uncorrected and highly corrected leads. *Circulation* 34: 638 1966
- Brody D A and Arzbaecher R C. The sampling and analysis of lead fields in electrocardiographic torso models. *IEEE Trans Bio Med Eng BME* 14:22 1967
- Lawrie T D V and Macfarlane P W. A clinical comparison of the Brody and Arzbaecher Frank McFee and Parungao corrected orthogonal lead systems. in Wenger R editor. *Actuelle Probleme der Vektorkardiographie VIII Internationales Kolloquium fur Vektorkardiographie* Vienna 1967. Published in Stuttgart 1968. Georg Thieme Verlag pp 110-118
- Macfarlane P W. A modified axial lead system for orthogonal lead electrocardiography. *Cardiovasc Res* 3:510 1969
- MacFarlane P W, Lorimer A R and Lawrie T D V. Normal ranges of modified axial lead system electrocardiographic parameters. *Br Heart J* 33: 258 1971
- Macfarlane P W, Lorimer A R and Lawrie T D V. Three and 12 lead electrocardiogram interpretation by computer. A comparison on 1093 patients. *Br Heart J* 33: 266 1971
- Gamboja R. Applicability of the axial lead system to infants and children. *Am J Cardiol* 18:690 1966
- Gamboja R and White N. The corrected orthogonal electrocardiogram in normal children. *McFee and Parungao System*. *AM HEART J* 75: 449 1968
- Ainger L E. Vectorcardiographic studies in infants and children. *Am J Cardiol* 21: 196 1968
- Ainger L E. Digital computer analysis of the vectorcardiogram of the newborn infant. Quantitative and comparative measurement of three orthogonal lead systems. *Circulation* 36:906 1967
- Witham A C. Quantitation of the vectorcardiogram. *AM HEART J* 72: 284 1966
- Penaloza D and Tranchesi J. The three main vectors of the ventricular activation process in the normal human heart. Its significance. *AM HEART J* 49:51 1955
- Sodi Palares D, Bisteni A, Medrano G A and Ayala C. *Clinical cardiopulmonary physiology* ed 2. New York 1960. Grune & Stratton Inc. p 63
- McCall B W, Wallace A G and Estes F H. Characteristics of the normal vectorcardiogram recorded with the Frank lead system. *Am J Cardiol* 10:514 1962
- Downs T and Liebman J. Statistical methods for vectorcardiographic directions. *IEEE Trans Bio Med Eng BME* 16:87 1969
- Brody, D A, Cox J W, McEachran A B, Giles H H and Ruesta V J. Spatial parameters and shape factors of the normal atrial vectorcardiogram and its scalar components. *Circulation* 39:229 1969
- Chou T and Helm R A. *Clinical vectorcardiography*. New York and London 1967. Grune & Stratton Inc.
- Snedecor G W. *Statistical methods* ed 5. Ames Iowa 1956. Iowa State University Press

Table III Comparison of surgical aspects of surviving and nonsurviving patients with coronary artery bypass grafts (55 patients)

| | Total no of grafts | Ave no grafts per pt | Removal of Lt wall | Isenberg procedure | Valvular surgery |
|-------------------|---|----------------------|---|--------------------|--|
| Survivors (44) | Saphenous vein 68 Internal mammary 9 Total 77 | 1.5 | Resection of wall 7 Plication of aneurysm 1 Total 8 | 1 | Plicate mitral valve 1 Replace mitral and aortic valves 1 Total 2 |
| Nonsurvivors (11) | Saphenous vein 19 Internal mammary 1 Total 20 | 1.8 | | 0 | Replace mitral valve 0 Mitral commissurotomy 1 Replace aortic valve 1 Total 2 |

Table IV Major complications associated with coronary artery bypass grafts (49 patients)

| Complication | Complication rate (no of patients) | % [No of pts with complications ÷ total (49 patients)] |
|--|---------------------------------------|---|
| Complications | | |
| Pulmonary emboli | 10 | 20 |
| Myocardial infarction | 9 | 18 |
| Brain damage | 9 | 18 |
| Postoperative hemorrhage | 7 | 14 |
| Cardiac arrest | 5 | 10 |
| Serum hepatitis | 3 | 6 |
| Renal failure | 2 | 4 |
| Subtotal | 45 | |
| Miscellaneous major complications | | |
| Chronic heart failure | 2 | |
| Tracheostomy | 2 | |
| Permanent pacemaker | 1 | |
| Postperfusion syndrome | 1 | |
| Gastrostomy | 1 | |
| Wound infection (death) | 1 | |
| Pneumonia | 1 | |
| Ulnar neuropathy (requiring surgery) | 1 | |
| Subtotal | 10 | |
| Total number of major complications | 55 | |

there were 35 major complications and six deaths. Over the last 15 month period 21 patients had CABG surgery and there were 20 major complications and five deaths.

Angiograms were done two to three weeks postoperatively on 27 of the 44 patients who survived the operations. Table VI shows that 71 per cent of the saphenous vein and 78 per cent of the internal mammary grafts were patent.

The over all patency rate was 44 out of 61 or 72 per cent.

The data suggest that a myocardial infarction may be associated with the CABG procedure even though one or more grafts remain patent. Five patients who had sustained an operative myocardial infarction were studied by angiography. Three of these patients each had received a single graft and in two the graft was patent.

Table I Comparison of clinical characteristics of surviving and nonsurviving patients with coronary artery bypass grafts (55 patients)

| | No of patients | Sex M/F | Age (yr) | Clinical class | | | | | | History of* | | | | |
|--------------|----------------|---------|----------|----------------|---|----|----|---|---|---------------|--------------|----------------|-------------------|-------------|
| | | | | 1 | 2 | 3 | 4 | 5 | 6 | Heart failure | Hypertension | Hyperlipidemia | Diabetes mellitus | Previous MI |
| Survivors | 44 | 35/9 | 53 | 5 | 6 | 12 | 19 | 2 | 3 | 6/41 | 11/42 | 10/42 | 5/42 | 22/43 |
| Nonsurvivors | 11 | 7/4 | 63 | 1 | 0 | 2 | 7 | 1 | 3 | 5/10 | 4/10 | 2/10 | 2/10 | 5/9 |

*Ratio of patients with positive history to total number of patients with known history

Table II Comparison of angiographic features of surviving and nonsurviving patients with coronary artery bypass grafts (55 patients)

| | Greater than 50% narrowing of major coronary artery* | | Abnormal LV contraction | Discrete LV aneurysm | Mitral insufficiency |
|-------------------|--|----------------|-------------------------|----------------------|----------------------|
| | No of involved arteries | No of patients | | | |
| Survivors (44) | 1 2 3 | 7 24 13 | 29 | 6 | 14 |
| Nonsurvivors (11) | 1 2 3 | 2 2 7 | 6 | 3 | 5 |

*Right coronary artery left circumflex coronary artery left anterior descending coronary artery

significance. One of the survivors had an internal mammary artery implanted (Vineberg operation) as an additional procedure. Noncontracting areas of the left ventricular wall were removed in eight of the survivors but in none of the nonsurvivors. Valvular surgery was also done in two of the survivors and in four of the nonsurvivors.

Table IV summarizes what were considered to be the major postoperative complications excluding death. Six patients died on the operating table. Fifty-five major complications were found in the remaining 49 patients. In this group 15 patients had one major complication, six had two, five had three, two had four, and one patient had five complications. There were five postoperative deaths. Of the 44 patients who survived, 20 had no major complication.

The major complications most frequently

recognized were pulmonary emboli (ten patients), myocardial infarction (nine patients), severe brain damage (nine patients), severe postoperative hemorrhage (five of the seven patients required reoperation), and cardiac arrest (five patients). As classified in Table IV, there were 10 miscellaneous major complications. It was of interest that two patients had no cardiac failure before operation but suffered from chronic congestive failure postoperatively.

Complications considered as minor are summarized in Table V. There was a total of 28 patients who had 17 different minor complications. Nine of the 44 patients who survived had neither a major nor minor complication.

The 55 patients were operated upon over a 30-month interval. In the first half of this period 34 patients had operations and

urgery and to a lesser degree a history of systemic hypertension. However a history of hyperlipidemia, diabetes mellitus or previous myocardial infarction was not more frequent in the nonsurvivors than the survivors.

Table III shows that details of the operation did not appear to distinguish the CABG survivors from the nonsurvivors. The average number of grafts done per patient was not statistically different in the two groups.

Major postoperative complications were frequent as summarized in Table IV. The most frequently recognized major complications included pulmonary emboli, myocardial infarction, severe brain damage, major postoperative hemorrhage and cardiac arrest. Table V shows that a wide variety of what were classified as minor complications also occurred. Nine of the 44 patients who survived had neither a major nor a minor complication.

The complications which are tabulated were those observed during regular clinical care. It is probable that a comprehensive search in the patients would have revealed other complications. For example, although pulmonary emboli were frequently diagnosed, the incidence might have been greater had each patient been specifically studied.

Both the mortality rate and the frequency of major complications were essentially unchanged for the first and last 15 months of the study. In this series increasing surgical experience did not appear up to this time to reduce the immediate risks. However it is possible that continued experience might produce improved results.

Table VI shows that the average early graft patency rate was 72 per cent. There appeared to be no particular difference for patency rate between the saphenous vein and the internal mammary grafts, although this is not certain because the latter group was small in number. It appeared possible that the placement of a CABG might be associated with accelerated closure of a stenotic coronary artery. Others have also reported this finding.² The patency of the graft apparently did not provide protection against a myocardial infarction complicating the operation.

Table VI Results of early postoperative angiograms of patients with coronary artery bypass grafts (27 patients)

| Type of graft | No of grafts done | No of grafts patent | Per cent of grafts patent |
|------------------|-------------------|---------------------|---------------------------|
| Saphenous vein | 57 | 37 | 71 |
| Internal mammary | 9 | 11 | 78 |
| Total | 61 | 44 | 72 |

Although the tables statistically summarize the results observed in the patients undergoing CABG, they do not adequately solve the physician's problem when he is faced with the decision of medical vs surgical treatment. As far as the immediate surgical procedure is concerned, the practicing physician wishes to know not only his patient's chances of survival but also the postoperative hazards. The following analysis illustrates one approach to arriving at a clinician's overview of the operation for this series of patients.

If we assume that the value of the CABG is to supply otherwise unavailable blood to the myocardium, in this series the operation was an immediate failure for those patients who died (11 patients) plus those known to have no patent graft (five patients). Adding from the remaining 39 patients the seven who had a myocardial infarction associated with the operation gives a cumulative failure rate of 23/55 or 42 per cent. Assuming that all the remaining 32 patients will have patent grafts and that all will achieve the full theoretical value of the operation, they must still run the gauntlet of major complications. Three patients had severe brain damage and of the remaining 29, two had cardiac arrest. Of the remaining 27, two had severe postoperative hemorrhage, of the 25, four had pulmonary emboli, of the 21, one had serum hepatitis. Thus the patient submitted for CABG had about two chances out of three of either dying, having no patent graft, or having a life-threatening postoperative complication.

Although the experience with the CABG in this series seems relatively clear, there

Table V *Minor complications associated with coronary artery bypass grafts (49 patients)*

| Complication | Complication rate (no of patients) |
|---|---------------------------------------|
| Femoral vein or artery thrombosis (requiring surgery) | 2 |
| Heart failure | 2 |
| Wound infection | 2 |
| Wound dehiscence | 2 |
| Phlebotrombosis | 2 |
| Hematoma of groin (requiring surgical drainage) | 2 |
| Respiratory insufficiency (requiring reintubation) | 2 |
| Urinary tract infection | 1 |
| Pericarditis | 1 |
| Postpericardiotomy syndrome | 1 |
| Ileus | 1 |
| Nonunion of sternum (requiring surgery) | 1 |
| Acute depression | 1 |
| Fever of unknown origin | 1 |
| Atelectasis and pleural effusion | 1 |
| Arrhythmias (requiring treatment) | |
| Sinus arrest with syncope | 1 |
| Atrial fibrillation and flutter (producing hypotension) | 2 |
| Premature ventricular contractions (frequent runs) | 1 |
| Supraventricular tachycardia and arrhythmias | 1 |
| Ulnar neuropathy | 1 |
| Total | 28 |

Three grafts had been placed in the fourth patient and two were open. In the remaining case three of four grafts were patent yet the patient had had a large inferior and anteroapical myocardial infarction followed by chronic severe cardiac failure.

Postoperative angiograms provided some evidence to suggest that progressive narrowing might occur in a stenotic coronary artery which was grafted. This was noted in two patients mentioned above who had had a myocardial infarction associated with a single graft, the graft later being shown to be patent. Preoperatively the degree of narrowing of the right coronary artery in these two patients was 80 and 95 per cent. Postoperatively the artery was completely occluded in both cases. In one additional

patient the preoperative films showed the lumen of the left circumflex branch reduced to 60 per cent and that of the right coronary artery reduced to 90 per cent. Postoperative angiograms demonstrated that both vessels were completely occluded but in this patient the status of the graft could not be determined.

Discussion

The coronary artery bypass graft (CABG) is currently a commonly performed procedure.¹ Much of the information which has been reported has come from a relatively limited number of institutions where the operation has been developed and most of the experience obtained. In attempting to assess the future role of the operation it appears desirable to obtain as broad a base of information as possible. This would not only be of value in accumulating the surgical results, but would also assist in evaluating the possible extension of the operation to the community hospital.

The patients referred to in this report were cared for in a 365 bed private community hospital. Surgery was performed by two different groups of surgeons. Each surgical team has been doing a wide variety of cardiovascular and open heart surgery in this hospital for the past seven years.

The number of patients in this report is relatively small in absolute terms. The patients, however, were drawn from a population receiving medical care at the community level and were all seen in consultations by the author during their hospitalization. The series represents therefore a significant segment of the practice of the individual physician caring for cardiac patients.

A total of 55 patients underwent CABG. The hospital mortality rate was 20 per cent. Table I compares the group who survived with the group who did not. There was a heavy predominance of men in each group. The survivors had an average age ten years less than that of the nonsurvivors. The average clinical class was not statistically different for the nonsurvivors compared to the survivors.

A history of cardiac failure was much more frequent in those who did not survive

Renal abnormalities discovered by routine postangiography abdominal films

Arturo Meneses Major MC

Richard J. McCarty Lieutenant Colonel MC

Stephen P. Glasser Major MC*

El Paso, Texas

In 1955 Richards and associates¹ reported on the greater incidence of congenital genitourinary anomalies in patients with congenital heart disease. In view of this Abrams and Kaplan² pointed out that a film of the abdomen should be obtained 10 to 15 minutes after the injection of contrast media in all cases of congenital heart disease subjected to angiocardiology.³ The excellent urographic study thus obtained frequently demonstrates anomalies of the urinary tract.

Yet despite the 10% incidence of some renal or ureteral anomaly found at autopsy in the general population⁴ to our knowledge it has not been routine practice by everyone to obtain abdominal films post angiography in adult patients being evaluated for acquired heart disease. It is therefore our feeling that this point merits reemphasis. In fifty six consecutive cases of patients having either pulmonary or coronary angiography three cases were discovered to have significant renal pathology on postangiography abdominal films and are the subject of this report.

Case reports

Case 1 H. S. was a 38 year old man who had a several year history of exertional chest pain. Slight hypertension (140/90) was noted on physical examination which was otherwise normal. Laboratory data revealed a Type IV hyperlipoproteinemia. ECG and stress tests were normal. Coronary angiography revealed 50 to 75 per cent occlusion of three vessels. His postangiographic abdominal flat plate showed changes consistent with either unilateral intrarenal hydronephrosis or chronic pyelonephritis (Fig. 1).

Case 2 C. M. was a 30 year old woman who was admitted with sudden pleuritic chest pain and shortness of breath. She was taking oral contraceptives for birth control. Physical examination revealed an accentuated pulmonary component to the second heart sound. Lung scan with ventilation perfusion studies suggested a right lower lobe defect and pulmonary angiography confirmed occlusive lesions of several right pulmonary artery branches. A postangiographic abdominal film (Fig. 2) revealed clubbing of the superior calyx with loss of architecture suggesting focal pyelonephritis. Tuberculous nephritis was subsequently ruled out.

Case 3 R. F. was a 35 year old woman with a two-year history of nonexertional chest pain precipitated by anxiety. Physical examination and laboratory data were normal. Resting ECG was normal but a prior Master's Test was reported as positive. A repeat submaximal treadmill test was negative. Coronary angiography was entirely normal. The postangiographic abdominal film (Fig. 3)

* From the Cardiology Service, Department of Medicine, William Beaumont General Hospital, El Paso, Texas.
Received for publication Oct. 6, 1973.

Reprint requests to Stephen P. Glasser, M.D., Cardiology Service, LSU Medical Center, Box 3432, New Orleans, Louisiana 70113.

Presented at the Annual Meeting of the Society of Medical Editors, New Orleans, Louisiana, September 1973.

no adequate control group with which to compare the figures. The desirability and critical need for the availability of a control group in order to allow statistical analysis of treated and untreated groups has been emphasized.⁴

The results of this series suggest that there may be important limitations to the CABG procedure. The evaluation of the operation as reported has been largely optimistic.⁵⁻⁸ It would appear desirable to have available the results from a cross section of the various institutions now performing this type of surgery, including those at the community level. The risks of the procedure in its various settings could then be better gauged and the clinician would gain a more factual overview of the operation.

Summary

The early results of 55 patients who underwent the coronary artery bypass graft (CABG) procedure in a large private hospital are reviewed. The hospital mortality rate was 20 per cent. Excluding death, 55 major complications were found in the total group of patients. Fourteen patients had more than one complication. Twenty of the 44 surviving patients had no major complication. Early postoperative angiograms showed an average graft patency rate of 72 per cent. There was some evidence of accelerated closure occurring in a stenotic coronary artery which was grafted. From the physician's point of view the patient submitted to the CABG procedure had about two chances out of three of either dying, having no patent graft, or

having a life threatening postoperative complication.

The experience with these patients suggests that there may be significant limitations to the performance of this operation or, at least to its possible application in the community hospitals. It is hoped that we will acquire a broader base of knowledge about the CABG as it is performed in various institutions across the country.

The author gratefully acknowledges the valuable assistance of Mrs Jeanne Heinz, R.N., Cardiology Nurse, in the preparation of this report.

REFERENCES

1. Friedberg C K. Caution and coronary artery surgery (editorial). *Circulation* 45:727 1972.
2. Burch G E. Coronary artery surgery saphenous vein bypass (editorial). *Am Heart J* 82:137 1971.
3. Aldridge H E and Trimble A S. Progression of proximal coronary artery lesions to total occlusion after aortocoronary saphenous vein bypass grafting. *J Thorac Cardiovasc Surg* 62:7 1971.
4. Spodick D H. Revascularization of the heart—numerator in search of denominator (editorial). *Am Heart J* 81:149 1971.
5. Morris G C Jr, Reul G J, Howell J F, Crawford E S, Chapman D W, Beasley H L, Winters W L, Peterson P K, and Lewis J M. Follow up results of distal coronary artery bypass for ischemic heart disease. *Am J Cardiol* 29:180 1972.
6. Johnson W D and Lepley D Jr. An aggressive surgical approach to coronary disease. *J Thorac Cardiovasc Surg* 59:128 1970.
7. Spencer F C, Green G E, Tice D A, and Glassman E. Bypass grafting for occlusive disease of the coronary arteries. A report of experience with 195 patients. *Ann Surg* 173:1029 1971.
8. Favaloro R G. Surgical treatment of coronary atherosclerosis by the saphenous vein graft technique. Critical analysis. *Am J Cardiol* 28:493 1971.

Renal abnormalities discovered by routine postangiography abdominal films

Arturo Venesio Major MC

Richard J. McCarty Lieutenant Colonel MC

Stephen P. Glasser Major MC*

El Paso Texas

In 1955 Richards and associates¹ reported on the greater incidence of congenital genitourinary anomalies in patients with congenital heart disease. In view of this, Abrams and Kaplan pointed out that a film of the abdomen should be obtained 10 or 15 minutes after the injection of contrast media in all cases of congenital heart disease subjected to angiocardiology.² The excellent urographic study thus obtained frequently demonstrates anomalies of the urinary tract.

Yet despite the 10% incidence of some renal or ureteral anomaly found at autopsy in the general population³ to our knowledge it has not been routine practice by everyone to obtain abdominal films postangiography in adult patients being evaluated for acquired heart disease. It is therefore our feeling that this point merits reemphasis. In fifty six consecutive cases of patients having either pulmonary or coronary angiography three cases were discovered to have significant renal pathology on postangiography abdominal films and are the subject of this report.

Case reports

Case 1. H. S. was a 38 year old man who had a several year history of exertional chest pain. Slight hypertension (140/90) was noted on physical examination which was otherwise normal. Laboratory data revealed a Type IV hyperlipoproteinemia. ECG and stress tests were normal. Coronary angiography revealed 50 to 75 per cent occlusion of three vessels. His postangiographic abdominal flat plate showed changes consistent with either unilateral intrarenal hydronephrosis or chronic pyelonephritis (Fig. 1).

Case 2. C. W. was a 30-year old woman who was admitted with sudden pleuritic chest pain and shortness of breath. She was taking oral contraceptives for birth control. Physical examination revealed an accentuated pulmonary component to the second heart sound. Lung scan with ventilation-perfusion studies suggested a right lower lobe defect and pulmonary angiography confirmed occlusive lesions of several right pulmonary artery branches. A postangiographic abdominal film (Fig. 2) revealed clubbing of the superior calyx with loss of architecture suggesting focal pyelonephritis. Tuberculous nephritis was subsequently ruled out.

Case 3. M. F. was a 35 year old woman with a two-year history of nonexertional chest pain precipitated by anxiety. Physical examination and laboratory data were normal. Resting ECG was normal but a prior Master's Test was reported as positive. A repeat submaximal treadmill test was negative. Coronary angiography was entirely normal. The postangiographic abdominal film (Fig. 3)

From the Cardiology Service, Department of Medicine, University of Texas Medical Branch at El Paso, El Paso, Texas. Received for publication October 4, 1972.

Reprint requests to Stephen P. Glasser, M.D., Cardiology Service, LSU Medical Center, Box 3332, Shreveport, Louisiana 71130.

Present address: Assistant Professor of Medicine, Louisiana State University Medical Center, Shreveport, Louisiana 71130.

is no adequate control group with which to compare the figures. The desirability and critical need for the availability of a control group in order to allow statistical analysis of treated and untreated groups has been emphasized.⁴

The results of this series suggest that there may be important limitations to the CABG procedure. The evaluation of the operation as reported has been largely optimistic.^{5,6} It would appear desirable to have available the results from a cross section of the various institutions now performing this type of surgery including those at the community level. The risks of the procedure in its various settings could then be better gauged, and the clinician would gain a more factual overview of the operation.

Summary

The early results of 55 patients who underwent the coronary artery bypass graft (CABG) procedure in a large private hospital are reviewed. The hospital mortality rate was 20 per cent. Excluding death, 55 major complications were found in the total group of patients. Fourteen patients had more than one complication. Twenty of the 44 surviving patients had no major complication. Early postoperative angiograms showed an average graft patency rate of 72 per cent. There was some evidence of accelerated closure occurring in a stenotic coronary artery which was grafted. From the physician's point of view the patient submitted to the CABG procedure had about two chances out of three of either dying, having no patent graft or

having a life threatening postoperative complication.

The experience with these patients suggests that there may be significant limitations to the performance of this operation, or, at least, to its possible application in the community hospitals. It is hoped that we will require a broader base of knowledge about the CABG as it is performed in various institutions across the country.

The author gratefully acknowledges the valuable assistance of Mrs. Jeanne Heinz, R.N., Cardiology Nurse, in the preparation of this report.

REFERENCES

1. Friedberg C K. Caution and coronary artery surgery (editorial). *Circulation* 45:727 1972.
2. Burch G E. Coronary artery surgery: saphenous vein bypass (editorial). *AM HEART J* 82:137 1971.
3. Aldridge H E and Trimble A S. Progression of proximal coronary artery lesions to total occlusion after aortocoronary saphenous vein bypass grafting. *J Thorac Cardiovasc Surg* 62:7 1971.
4. Spodick D H. Revascularization of the heart—numerators in search of denominators (editorial). *AM HEART J* 81:149 1971.
5. Morris G C Jr, Reul G J, Howell J F, Crawford E S, Chapman D W, Beazley H L, Winters W L, Peterson P K, and Lewis J M. Follow up results of distal coronary artery bypass for ischemic heart disease. *Am J Cardiol* 29:180 1972.
6. Johnson W D and Lepley D Jr. An aggressive surgical approach to coronary disease. *J Thorac Cardiovasc Surg* 59:128 1970.
7. Spencer F C, Green G E, Tice D A, and Glassman E. Bypass grafting for occlusive disease of the coronary arteries. A report of experience with 195 patients. *Ann Surg* 173:1079 1971.
8. Favaloro R G. Surgical treatment of coronary arteriosclerosis by the saphenous vein graft technique. Critical analysis. *Am J Cardiol* 28:193 1971.



Fig 2 Case No 2 The postangiography abdominal film reveals clubbing of the superior calyx of the right renal pelvis with loss of architecture suggesting focal pyelonephritis.

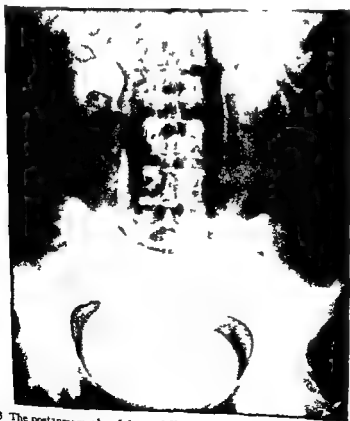


Fig 3 Case No 3 The postangiography abdominal film demonstrates segmental megaloureter and pelvic distension on the right.



Fig 1 Case No 1 The postangiography abdominal film demonstrates a right renal abnormality suggesting intrarenal hydronephrosis or unilateral chronic pyelonephritis

revealed segmental megalooureter and pelvic distension suggesting a compensated ureteropelvic junction obstruction

Discussion

The high incidence of genitourinary anomalies in the general adult population³ led us to obtain a routine abdominal film at the termination of the procedure in all patients who had undergone an angiographic study. Because high quality films can be obtained this would serve as an excellent screening test in patients with latent genitourinary disease.

In fifty six consecutive cases patients had postangiography abdominal films performed and three significant abnormalities were found. In one case clubbing of the right upper pole of the calyceal system with loss of architecture suggested focal pyelonephritis. Tuberculous nephritis was

sought with multiple urine cultures for acid fast bacilli but proved negative. In another case the urographic appearance was that of either chronic pyelonephritis or intrarenal hydronephrosis with spontaneous relief of a prior clinically asymptomatic obstruction. In the third instance segmental megalooureter and distension of the renal pelvis suggested a compensated ureteropelvic junction obstruction. (A fourth case not shown, demonstrated indentation of the ureter by an aberrant vessel but was of no clinical consequence.)

The results of this study lend support to our recommendation to obtain postangiographic abdominal films in all patients undergoing an angiographic procedure. The only risk is the addition of a small amount of radiation exposure which is not felt to be significant when compared to the information obtained.

Relation of microcirculatory thrombosis to thrombus in the proximal coronary artery Effect of aspirin, dipyridamole, and thrombolysis*

Christos B Moschos MD

Kamalesh Lahiri MD

Michael Lyons MD

Allen B Weisse MD

Henry A Oldewurtel

Timothy J Regan MD

Newark N J

Studies on coronary thrombosis have been traditionally focused on the primary thrombotic obstruction of a major epicardial vessel and the factors favoring its pathogenesis. While results derived from such studies remain of critical importance to the understanding of the determinants favoring the occurrence of coronary occlusion, other aspects of this process have received little or no attention. Sporadic observations made at autopsy have described the presence of thrombi or platelet aggregates in the intramyocardial vessels posing the question of their potential pathogenic role in patients with coronary artery disease^{1,2} or the dysfunction of other organ.^{3,4} However, the role of platelet aggregates in the small vessels distal to a thrombus in a major artery has received little consideration. Conceivably obstruction of myocardial microcirculation could

compromise effective collateral flow and thus intensify the effects of ischemia upon the myocardial area affected by proximal coronary occlusion. In previous studies we observed a substantial decline in the platelets of blood sampled sequentially from the coronary sinus during the formation of a thrombus in the coronary artery.^{5,7} We felt that the extent of the drop in platelets could not be solely due to thrombus formation in the epicardial coronary vessel. Since platelet aggregation is considered to be the important factor responsible for the initiation and extension of arterial thrombus,^{8,9} a study was designed to determine platelet distribution in the microcirculation of the myocardium after experimental thrombosis of a major coronary vessel. In addition, the recent studies suggesting a potential antithrombotic effect of aspirin¹⁰ and dipyridamole^{12,14} two of several agents with anti-

From the Department of Medicine, College of Medicine and Dentistry of New Jersey, Newark, N.J.

This study was supported in part by United States Public Health Service Research Grant HL 15050 and grants from the East County Heart Association of New Jersey.

Received for publication August 2, 1972.

Reprint requests to Christos B Moschos, MD, College of Medicine and Dentistry of New Jersey, 100 Bergen Street, Newark, N.J. 07103.

Presented in part at the 64th Meeting of The American Heart Association, Anaheim, California, 1971.

Summary

Patients with acquired heart disease undergoing angiographic procedures had routine postangiographic abdominal films performed. In fifty six consecutive cases with clinically unsuspected renal disease three patients were found to have significant abnormalities.

The not infrequent finding of clinically unsuspected renal pathology in patients undergoing angiographic procedures during the evaluation of acquired heart disease has led us to the recommendation that all patients have postangiographic abdominal films. This represents an inexpensive screening test for renal pathology with no increase in patient risk.

The authors wish to express their appreciation to the personnel of the Department of Medical Research and Development for their support in the completion of this manuscript.

REFERENCES

- 1 Richards M R, Merritt K K, Samuels M H and Longmann A G. Congenital malformations of the cardiovascular system in a series of 6053 infants. *Pediatrics* 15:12 1955
- 2 Abrams H L and Kaplan H S. *Angiographic interpretation in congenital heart disease*. Springfield 1956 Charles C Thomas Publisher p 14
- 3 Campbell M. *Textbook of urology*. Philadelphia 1954 W B Saunders Company p 227

thrombus) after thrombus formation¹⁰ The length of Thrombolytic infusion was two hours for the early thrombus (five dogs) and four hours for the late thrombus (six dogs) Group IV of 17 and Group V of 7 animals were given respectively 500 mg of aspirin and 100 mg of dipyridamole daily for seven days prior to induction of thrombus The animals were killed and autopsy was performed two to four hours after thrombus formation or immediately after spontaneous ventricular fibrillation The heart was excised the location of the thrombus was noted and its weight was determined Multiple tissue samples of comparable weight were taken from the ischemic and nonischemic area of the heart (Fig 1) from all or several dogs of each group for radioactivity counting and histologic examination (Table I) The radioactivity of the tissue was determined in a well type scintillation counter and expressed as counts per minute per Gm of tissue and also as a count ratio of the ischemic to nonischemic area of each dog after the individual readings of each area were averaged Histologic examination of the cardiac tissue was made in multiple sections stained by hematoxylin-eosin and phosphotungstic acid hematoxylin For statistical evaluation the results of each treated group were compared to the non treated thrombus group The formula for non paired observation was used for evaluation of thrombus weight and radioactivity ratios whereas for the incidence of arrhythmias the chi square formula was applied

Results

A thrombus was formed in 47 out of 56 animals in which it was attempted In Group III lysis of thrombus in the major coronary artery was complete in four and incomplete in three dogs In the remaining four dogs the thrombus did not lyse In Groups IV and V aspirin or dipyridamole failed to affect significantly the incidence or weight of the formed thrombus (Table II) Fig 1 shows schematically the distribution of radioactivity in animals without thrombus production In the three areas from which tissue samples were taken the radioactivity of the labelled platelets was quite comparable at approximately 50

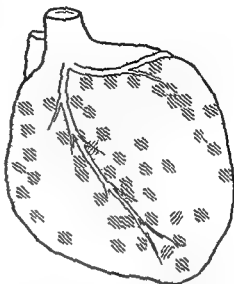


Fig 1 Schematic illustration of distribution of radioactivity in control animals infused with ^{51}Cr platelet without induction of thrombus The small squares represent the locations from which the tissue samples were taken for radioactivity counting Samples for histology were taken from adjacent areas The shaded part represents schematically the intensity of radioactivity indicating uniform distribution in all segments

counts per minute per Gm of tissue In contrast following thrombus formation there were distinct areas of radioactivity (Fig 2) Tissue samples from nonobstructed regions presented low activity similar to that found in the no thrombus group An area of high activity was seen in the entire ischemic area distal to the occluding coronary thrombus The counts here obtained from tissue samples of weight comparable to those obtained from nonischemic areas ranged from 130 to 170 per minute per Gm The area adjacent to the main thrombus showed a much higher activity with counts 450 to 600 per minute per Gm of tissue while the thrombus itself showed 5 000 to 13 000 counts per minute per Gm of weight The radioactivity ratio of ischemic to control area when averaging the activity of all samples was 3.31 with a range of 2.8 to 9.5 (Table III) The ratio of thrombus radioactivity to that of the control area was 120 average with a range of 60 to 250 On histologic examination the typical findings in all sections from the ischemic myocardial tissue of the nontreated animals

Table I Groups studied

| Group | Control no thrombus | Thrombus no treatment | Thrombus + fibrinolysin | Aspirin + thrombus | Persantin + thrombus |
|--------------------|---------------------------|-----------------------------|-------------------------------|--------------------------|----------------------------|
| | I | II | III | IV | V |
| Number of animals* | 4 (4) | 12 (8) | 11 (10) | 17 (7) | 7 (7) |

*Numbers in parentheses indicate animals having microcirculatory assessment by histology and radioactivity counts

platelet aggregation properties: prompted us to assess their effect, as well as that of thrombolysis upon this preparation

Material and methods

A total of 67 apparently healthy male mongrel dogs fasting for 18 hours and weighing from 19 to 24 kilograms were used. They were anesthetized with morphine sulfate (3 mg per kilogram of body weight intramuscularly) and sodium pentobarbital (20 mg per kilogram of body weight intravenously) and subsequently placed on a respiratory pump to maintain adequate ventilation. The left jugular vein and left carotid artery were exposed through small skin incisions. After an estimate of platelet adequacy in each dog, 170 to 180 ml of blood was taken by means of a polyethylene catheter placed in the jugular vein infusing subsequently equal amounts of isotonic saline to maintain blood volume. Platelet rich plasma (PRP) was obtained and the platelet button was then labelled with 300 μ c of sodium chromate (^{51}Cr) as described by Aas and Gardner and modified by Aster and Janle.¹³ The separation of platelets from canine blood was accomplished by repeated centrifugation of the initial blood sample at speeds ranging from 1 000 to 1 800 r p m for four to seven minutes. After obtaining PRP with 75 per cent of the original platelet count as a rule the plasma was centrifuged at high speeds to produce the platelet button. After tagging with ^{51}Cr the platelets were infused into the dog and 1 to 1½ hours were allowed for equilibration.

At this time production of a platelet coronary thrombus was attempted in 63 dogs by means of a catheter electrode

as previously described.^{8,16} In brief this method employs a No 8 Sones catheter containing a stainless steel insulated wire placed in the anterior descending or circumflex coronary artery. By means of a dry cell battery connected to the proximal end of the electrode wire, a 300 to 500 micro ampere current is applied to the intima of the vessel, resulting in platelet thrombus formation within 15 to 90 minutes. Morphologically the thrombus is similar to that described in human arteries.^{18,19} Its formation in this model is usually indicated by appearance of acute injury potential and multiple premature ventricular beats in the continuously monitored ECG. Appearance of injury potential within the first one to two minutes following the placement of the catheter is in our experience, due to mechanical obstruction of the coronary vessel by the catheter. Seven such animals were excluded from the study thus leaving a total of 56 dogs with attempted coronary thrombosis. The four remaining dogs of the initial group of 67 which received ^{51}Cr tagged platelets had a catheter placed in the coronary artery without attempting induction of thrombosis (Group I). According to the treatment of the animals either before thrombus induction or after thrombus formation the rest of the dogs were divided into various groups (Table I) the no treatment (Group II) comprised 12 animals which received no treatment prior to thrombus induction. Eleven animals (Group III) were infused into the left ventricle with Thrombolysin * 2,500 units per kilogram of body weight per hour one hour (early thrombus) or three hours (late

*Merck Sharp and Dohme West Point Pa

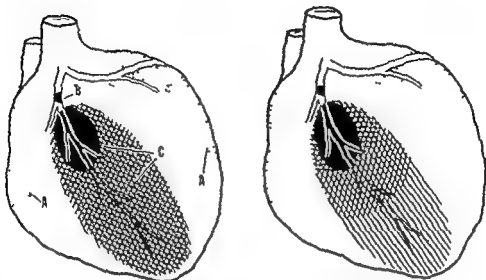


Fig. 2. Left panel: Schematic illustration of distinct area of radioactivity following thrombus formation. A = control area corresponding to nonobstructed coronary artery showing low activity. B = occluding thrombus. C = area of high activity which is more prominent in the region adjacent to the formed thrombus. Right panel: Distribution of radioactivity in animals pretreated with aspirin and dipyridamole. There is a definite decrease in radioactivity, becoming minimal in the distal half of the ischemic region.

they were not included in the table because the animals were given antiarrhythmics. This was necessary in order to have two to four hour survivors following thrombus formation which could subsequently be infused with fibrinolytic. In this preparation the mortality rate of the animals due to the presence of an obstructing coronary thrombus is approximately 50 per cent within the first two hours and about 90 per cent within four hours following formation of thrombus.²³

Discussion

The results of this study suggest that thrombosis in a major coronary artery is as occurred at least in the first hours following its occurrence with the presence of platelet thrombi in the microcirculation of the ischemic area. The mechanism of formation is not clear. Embolization of platelet aggregates from the primary site¹ represents a plausible pathogenetic route. Similar mechanisms have been presumed to operate in the microcirculation of other organs such as brain⁴ or kidney² resulting in damage of the involved tissue. Jørgensen and col-

leagues²³ have shown that platelet aggregates induced by adenosin diphosphate infusion into the coronary artery or in the aorta proximal to the renal arteries results in myocardial or renal injury respectively. James¹ has suggested that microvascular coronary occlusion by platelet aggregates may play a significant role in the pathogenesis of human myocardial necrosis.

In the present study microcirculatory thrombosis was consistently observed in association with major (epicardial) artery thrombosis. The role of microvascular thrombosis in the injury produced in ischemic myocardium is open to speculation. It is generally agreed that the important factor determining the extent of myocardial damage associated with coronary obstruction is the degree of availability of collateral blood flow. The present findings suggest that following proximal obstruction by a thrombus access of available collateral anastomotic flow distally into the ischemic area can be compromised by the presence of multiple platelet aggregates in the intramyocardial vessels although direct evidence for this view is not yet available. It

Table II Effect of aspirin and Persantin on major coronary artery thrombosis

| Group | Thrombus no treatment | Aspirin + thrombus | Persantin + thrombus |
|---------------------------------|--------------------------|-----------------------|-------------------------|
| Number of animals | 12 | 17* | 7 |
| Weight (mg) of thrombus and SE† | 63 ± 20 | 69 ± 33 | 114 ± 46 |

*No thrombus was formed in three animals
 †SE = standard error of the mean

Table III Effect of aspirin, dipyridamole, and Thrombolytic on average radioactivity ratios of ischemic to control areas (I/C)

| Group | Control no thrombus | Thrombus no treatment | Thrombus + Thrombolytic | Aspirin + thrombus | Persantin + thrombus |
|-------------------|---------------------------|-----------------------------|-------------------------------|--------------------------|----------------------------|
| | I | II | III | IV | V |
| Number of animals | 4 | 8 | 11 | 7 | 7 |
| I/C ratio | 1.0 | 3.30 | 1.84* | 1.35* | 2.01* |

*P < 0.01

was the presence of multiple randomly distributed microthrombi occluding the arterioles and capillaries of the ischemic area. These thrombi mainly consisted of platelets and minimal amounts of fibrin (Fig 3). In the area adjacent to the main thrombus a variable degree of thrombus propagation into the side branches could be observed.

Pretreatment of the animals with aspirin and dipyridamole or infusion with Thrombolytic, modified the patterns of radioactivity and histology of the microcirculation observed in the nontreated animals (Table III). There was reduced radioactivity in the ischemic area resulting in a significant ($P < 0.01$) decrease of the ischemic/nonischemic ratio in all treated groups; however the distribution pattern of radioactivity varied among Groups III, IV and V. In the dogs infused with Thrombolytic there was a more or less uniform radioactivity decrease in the entire ischemic region. This pattern remained the same whether or not the thrombus was totally, partially or not at all lysed (Table IV). The radioactivity distribution in the ani-

mals receiving aspirin or dipyridamole is shown schematically in Fig 2. The main decrease in radioactivity in these animals was observed in the distal half of the ischemic region whereas in the remaining area particularly adjacent to the thrombus the change was smaller. Regarding histologic findings in the animals treated with fibrinolytic there was virtually no evidence of microthrombi in sections from the ischemic myocardium. However it should be noted that in Groups IV and V disappearance of microthrombi followed a pattern similar to that described above for radioactivity. It was observed only in the sections taken from the distal ischemic myocardium whereas in many of the sections taken proximal to the thrombus some degree of microthrombosis was present.

There was a decrease in the incidence of arrhythmias and mortality from ventricular fibrillation in the treated animals although it was found significant only for the aspirin group (Table V). The incidence of arrhythmias and mortality in the animals receiving fibrinolysis was also significantly lower when compared to control; however

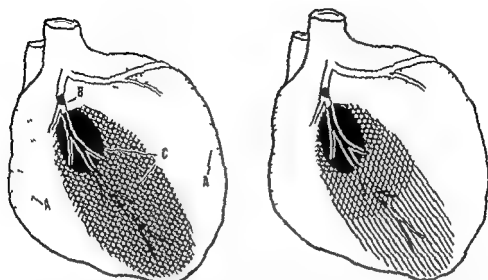


Fig. 2. Left panel: Schematic illustration of distinct areas of radioactivity following thrombus formation. A = control area corresponding to nonobstructed coronary artery showing low activity. B = occluding thrombus. C = area of high activity which is more prominent in the region adjacent to the formed thrombus. Right panel: Distribution of radioactivity in animal pretreated with aspirin and dipyridamole. There is a definite decrease in radioactivity becoming minimal in the distal half of the ischemic region.

they were not included in the table because the animals were given antiarrhythmics. This was necessary in order to have two to four hour survivors following thrombus formation which could subsequently be infused with fibrinolytic. In this preparation the mortality rate of the animals due to the presence of an obstructing coronary thrombus is approximately 90 per cent within the first two hours and about 90 per cent within four hours following formation of thrombus.¹⁶

Discussion

The results of this study suggest that thrombosis in a major coronary artery is associated at least in the first hours following its occurrence with the presence of platelet thrombi in the microcirculation of the ischemic area. The mechanism of formation is not clear. Embolization of platelet aggregates from the primary site represents a plausible pathogenetic route. Similar mechanisms have been presumed to operate in the microcirculation of other organs such as brain¹⁷ or kidney¹⁸ resulting in damage of the involved tissue. Jørgensen and col-

leagues^{19,20} have shown that platelet aggregates induced by adenosin diphosphate infusion into the coronary artery or in the aorta proximal to the renal arteries results in myocardial or renal injury respectively. James²¹ has suggested that microvascular coronary occlusion by platelet aggregates may play a significant role in the pathogenesis of human myocardial necrosis.

In the present study microcirculatory thrombosis was consistently observed in association with major (epicardial) artery thrombosis. The role of microvascular thrombosis in the injury produced in ischemic myocardium is open to speculation. It is generally agreed that the important factor determining the extent of myocardial damage associated with coronary obstruction is the degree of availability of collateral blood flow. The present findings suggest that following proximal obstruction by a thrombus access of available collateral anastomotic flow distally into the ischemic area can be compromised by the presence of multiple platelet aggregates in the intramyocardial vessels although direct evidence for this view is not yet available. It



Fig 3 Photomicrograph of capillary and arteriol (left) with small distended artery (right) all occupied by platelet thrombi (Hematoxylin and eosin. Original magnification $\times 250$)

Table IV Effect of lysis of main thrombus on radioactivity ratios of ischemic/control areas

| | Thrombus | Thrombus and Thrombolysis | | |
|--------|----------|---------------------------|---------------|-------------------|
| | | Complete lysis | Partial lysis | Occuding thrombus |
| Number | 8 | 4 | 3 | 4 |
| I/C | 3.30 | 1.82 | 1.89 | 1.80 |

was further shown that a relatively short infusion of a thrombolytic agent is capable of minimizing significantly the extent of microcirculatory thrombosis, an effect which was independent of the capability of the thrombolytic agent to dissolve the primary thrombus (Table IV). The same result was obtained following premedication for a relatively brief period with aspirin or persantin which, while failing to affect thrombosis in a major coronary artery as we have demonstrated previously,¹¹ modified platelet behavior in the microcirculation. Further analysis of these effects would indicate that while platelet aggregation seems to be the primary mechanism responsible for obstruction of the microcirculatory vessels, fibrin as shown in sections stained with phosphotungstic acid hematoxylin appear at some stage of the

process thus explaining the effect of the thrombolytic agent whose action is directed only against fibrin.

On the other hand the effects of aspirin and dipyridamole presumably due only to their antiplatelet aggregation properties should be considered more specific by virtue of their ability to prevent initial platelet aggregation. Regarding their effectiveness at the microcirculatory but not the epicardial vessel level it should be pointed out that while thrombolytic agents can be variably successful depending on the age of the thrombus,¹² the effectiveness of the aspirin and dipyridamole might be determined by other factors such as size of the vessel¹³ or the intensity of the aggregating stimulus in the proximal artery—the electric current. Presumably the levels of aspirin and dipyridamole in this study were

not adequate to protect against thrombus formation in vessels with diameters much larger than that of the microcirculation. Alternatively, the favorable results against thromboembolism shown by the studies of Salzman and colleagues¹⁴ and Sullivan and associates¹⁵ suggest that aspirin and dipyridamole are effective only where thromboembolism rather than thrombosis is involved. This in turn would indicate that in the pathogenesis of microcirculatory thromboembolism is an important mechanism. It is recognized that the occurrence of microthrombi in disease states might be determined by other variables which are not necessarily related to the conditions of the model used in this study.

Although the significance of microcirculatory thrombosis will have to be further assessed, the decrease in incidence of arrhythmias and mortality observed in the groups of animals pretreated with aspirin and dipyridamole is worth emphasizing. However, only in the aspirin group where the decrease of microcirculatory thrombosis was more impressive (Table III) was it found to be statistically significant (Table V). In this regard it is interesting to note that Hjerem¹⁶ found a higher number of intramyocardial arteries occupied by platelet aggregates in patients with known coronary artery disease who died suddenly as compared to those patients who died of noncardiac causes. Occlusion of small coronary vessels either supplying directly or responsible for anastomotic access to the various parts of the conduction system can potentially cause disturbances of the electrophysiologic function of the heart.¹

Our observations—that fibrinolytic agents aspirin or persantin while failing to affect thrombosis in the major coronary artery modify the platelet behavior in the microcirculation of this model—indicate the need for continuing studies in this direction. If collateral flow in an obstructed coronary arterial system is the factor of critical importance for electrical stability, survival and repair of the ischemic myocardium, it is conceivable that such pathology of the microcirculation might be responsible for compromising the effectiveness of this important compensatory mechanism. On the other hand, metabolic aspects relevant to cardiovascular function

Table V Incidence of arrhythmias and ventricular fibrillation

| Group | No. of animals | Arrhythmias | Ventricular fibrillation |
|-----------|----------------|-------------|--------------------------|
| Control | 12 | 10 (83%) | 8 (66.7%) |
| Aspirin | 17 | 5 (30%) | 2 (12%) |
| Persantin | 7 | 3 (43%) | 2 (29%) |

$P < 0.01$

†P = not significant

should be explored particularly in regard to aspirin whose variable effects upon metabolism have been described.¹¹

Summary

A study was designed to determine the distribution of platelets in the microcirculation of the myocardium following experimental platelet thrombosis in a major coronary vessel. Determination was made by means of ⁵¹Cr labelled platelets and histologic examination. The results suggest that thrombosis in a major coronary artery is associated at least in the first hours following its occurrence with the presence of platelet thrombi in the microcirculation of the ischemic area. Pretreatment with aspirin or dipyridamole minimized significantly the extent of microcirculatory thrombosis without affecting the thrombus in the proximal coronary artery. A thrombolytic agent given after thrombus formation had a similar effect upon microcirculatory thrombosis remaining also independent of its effect upon proximal coronary artery thrombosis. There was a decrease in incidence of arrhythmias and mortality rate in the group of animals pretreated with aspirin and dipyridamole.

The authors wish to thank Miss Elaine Podell and Mrs. Amparo Escobinas for technical assistance and Mrs. Audrey Brown for secretarial services rendered.

REFERENCES

1. James T. N. Pathology of small coronary arteries. *Am J Cardiol* 20:679, 1967.
2. Jørgensen L. The role of platelet embolism from crumbling thrombi and of platelet aggregates arising in flowing blood in Sherry S.

- Brinkhous K, Genton E and Stengle J editors Thrombosis Washington D C 1969 National Academy of Sciences pp 506 535
- 3 Russel R Observations on the retinal blood vessels in monocular blindness *Lancet* I 1422 1961
- 4 Meyer J S Gotoh T and Tazaki Y Circulation and metabolism following experimental cerebral embolism *J Neuropathol Exp Neurol* 21:4 1962
- 5 Moore S and Mercer W A Microembolic renal ischemia and hypertension *Crit Med Assoc J* 92 221 1965
- 6 Moschos C B Lehan P H Korovenidis G T Weiss A B Oldewurtel H A and Regan T J Regional and systemic clotting and fibrinolytic activity after myocardial infarction *Am J Physiol* 216 (Suppl 2) 308 1969
- 7 Moschos C B Oldewurtel H A and Regan T J Platelet counts during acute coronary thrombosis *Circulation* 42 (Suppl III) 185 1970
- 8 Glynn M F Murphy E A and Mustard J F Platelet and thrombosis *Ann Intern Med* 61 715 1966
- 9 Weiss H J and Aledort L M Impaired platelet/connective tissue reaction in man after aspirin ingestion *Lancet* II 495 1967
- 10 O'Brien J R Effects of salicylates on human platelets *Lancet* I:779 1966
- 11 Atac A Spagnuolo M and Zucker M H Long term inhibition of platelet functions by aspirin *Proc Soc Exp Biol Med* 133 1331 1970
- 12 Emmons P R Harrison M J G Honour A J and Mitchell J R A Effect of a pyrimidin derivative on thrombus formation in the rabbit *Nature* 208 255 1965
- 13 Didisheim P Inhibition by dipyridamole of arterial thrombosis in rats *Thromb Diath Haemorrh* 20:257 1968
- 14 Sullivan J M Harken D E and Gorlin R Pharmacologic control of thromboembolic complications of cardiac valve replacement *N Engl J Med* 279 576 1968
- 15 Aster R H and Jandl J H Platelet sequestration in man I Methods *J Clin Invest* 43 843 1964
- 16 Salazar A E Experimental myocardial infarction *Circ Res* 9 1351 1961
- 17 Constantinides P Plaque fissures in human coronary thrombosis *J Atheroscler Res* 61 1 1966
- 18 Moschos C B Lahiri K Peter A Jesrani M and Regan T J Effect of aspirin upon experimental coronary and noncoronary thrombosis and arrhythmias *Am Heart J* 81 575 1972
- 19 Moschos C B Burke W M Lehan P H Oldewurtel H A and Regan T J Thrombolytic agents and lysis of coronary artery thrombosis *Cardiovasc Res* 4 228 1970
- 20 Weiss A B Moschos C B Passanante A J Khan M I and Regan T J Relative effectiveness of three antiarrhythmic agents in the treatment of ventricular arrhythmias in experimental acute myocardial ischemia *Am Heart J* 81 503 1971
- 21 Bertram H J and Fulton G P Platelets in the peripheral circulation in Johnson S A Monto R W Rebeck J W and Horn R C Jr editors Blood platelets Henry Ford Hospital International Symposium Boston 1961 Little Brown & Company pp 7 19
- 22 Jørgensen L Rowsell H C Hovig T Glynn M F and Mustard J F Adenosine diphosphate induced platelet aggregation and myocardial infarction in swine *Lab Invest* 17 616 1967
- 23 Jørgensen L Glynn M F Hovig T Murphy E A Buchanan M R and Mustard J F Renal lesions and rise in blood pressure caused by adenosine diphosphate-induced platelet aggregation in rabbits *Lab Invest* 23 347 1970
- 24 Salzman E W Harris W H and DeSanctis R W Reduction in venous thromboembolism by agents affecting platelet function *N Engl J Med* 281:1287 1971
- 25 Sullivan J M Harken D E and Gorlin R Pharmacologic control of thromboembolic complications of cardiac valve replacement *N Engl J Med* 281 1391 1971
- 26 Haerem J W Platelet aggregates in intramyocardial vessels of patients dying suddenly and unexpectedly of coronary artery disease *Atherosclerosis* 15 199 1972
- 27 Smith M J H Metabolic effects of salicylates in Smith M J H and Smith P K editors The salicylates A critical review New York 1966 Interscience Publishers Inc p 49

Electrophysiological effects of lidocaine on distal Purkinje fibers of canine heart

John Witting MD

Lura A Harrison PhD

Andrew G Wallace MD*

Durham NC

The tertiary amine lidocaine hydrochloride was first noted to have antiarrhythmic effects on the hearts of experimental animals in 1950.¹ Subsequently it was shown that injections of lidocaine either abolished or markedly attenuated the ventricular premature beats which occurred in patients after myocardial infarction^{2,3} after open heart surgery^{4,5} and in situations of presumed digitalis overdose.^{6,7} In 1969 Sugimoto and associates⁸ reported certain of the electrophysiological effects of lidocaine on the intact heart and noted marked differences between the effects of lidocaine and either quinidine or procainamide. Davis and Temte⁹ noted that in isolated Purkinje fibers lidocaine shortened the duration of the action potential and failed to significantly reduce the maximal rate of rise of phase 0.

In a recent report Myerburg and colleagues¹⁰ called attention to a marked disparity between the times of repolarization in adjacent distal Purkinje fibers. The long duration of action potentials and refractoriness in some Purkinje fibers could prevent the propagation of premature beats

across the segment of distal Purkinje tissue with the longest action potential duration. This segment functioned as a gate. The gating function of distal Purkinje fibers was ascribed to longer action potentials and their long refractory period was thought to contribute to decremental conduction and block of early premature impulses. Because lidocaine has been shown to shorten action potential duration this study was undertaken to examine the effects of lidocaine on distal Purkinje fibers and the functional consequences of these effects on the propagation of early premature beats. The results of this study suggest a potentially important mechanism for the antiarrhythmic action of lidocaine and a model system for the study of other antiarrhythmic agents.

Methods

These experiments were performed on isolated cardiac tissues obtained from healthy adult mongrel dogs. The dogs were anesthetized with sodium pentobarbital 30 mg per kilogram of body weight given intravenously and the heart was removed

From the Department of Medicine and Physiology, Duke University Medical Center, Durham, NC. This work was supported in part by grants from the United States Public Health Service HL-05545 and HL-05736 and from the North Carolina Heart Association.
Received for publication August 4, 1972.
Reprint requests to Andrew G. Wallace, MD, Box 302, Duke University Medical Center, Durham, NC 27710.
Dr. Wallace is the recipient of a United States Public Health Service Research Career Development Award.

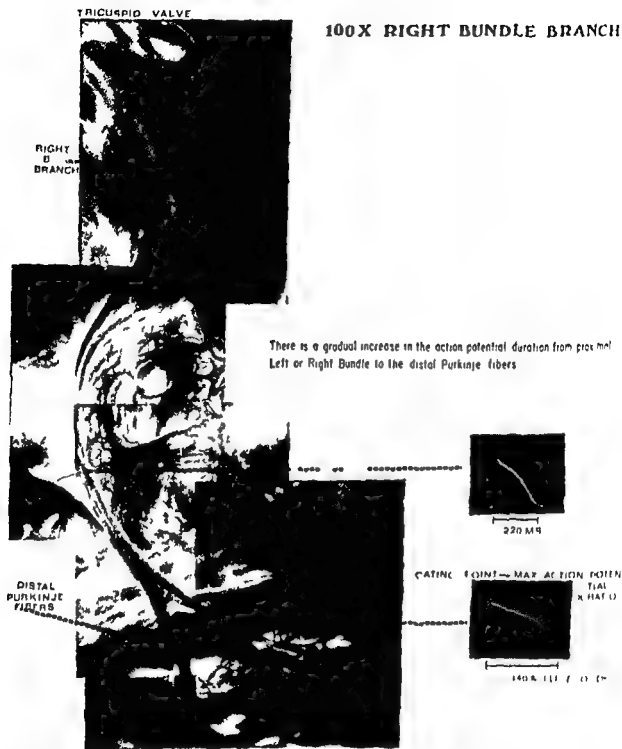


Fig 1 Photograph of the preparations used in these studies. The photo is of the right septal surface. The septal leaflet of the tricuspid valve is above. The anterior papillary muscle projects to the left in the center of the photo. The right bundle branch and free running distal Purkinje fibers have been stained dark with Lugol's solution. Typical action potentials from the proximal right bundle branch and distal Purkinje fibers are shown in the inserts.

through a right thoracotomy. Longitudinal cuts were made parallel to the ventricular septum, one along the anterior interventricular groove and one along the posterior interventricular groove. The chordae tendineae were then cut at their attachment to the tricuspid

valve and the right ventricular free wall was reflected away from the septum. The right bundle branch was identified at the level of the membranous septum and was transected at this level. A layer of septal muscle, 3 to 5 mm in thickness and containing

the right bundle branch was dissected and removed. At the base of the anterior papillary muscle the right bundle branch became a free running strand with two or more branches which inserted into the endocardial surface of the right ventricular free wall. A section of the free wall containing these insertions also was removed. The preparation thus included (1) the proximal right bundle branch (2) the anterior papillary muscle (3) the false tendon(s) and (4) an endocardial section of the right ventricle to which the false tendons inserted (see Fig. 1). After transferring this preparation to a tissue bath the muscle between the anterior papillary muscle and the right ventricular wall was divided so that the false tendon(s) was the only connection between the right bundle branch and the right ventricular wall. The tissue was then fixed to the elastic bottom of the bath with small stainless steel pins and perfused at a rate of 10 ml per minute with a modified Tyrode solution which was equilibrated with 95 per cent oxygen and 5 per cent carbon dioxide. The Tyrode solution was composed of the following (mM concentration): NaCl 137, NaHCO_3 12, CaCl_2 2.2, KCl 5.4, MgCl_2 0.5, NaH_2PO_4 1.8 and dextrose 5.0. Temperature within the bath was maintained at $38 \pm 0.5^\circ \text{C}$ by an electrofilm heating element beneath the silver bath. The film was powered by a fully rectified direct current source and was regulated by a thermocouple within the bath and a feedback control circuit.

Transmembrane action potentials were recorded with glass microelectrodes. The microelectrodes were machine pulled with a Kopf vertical pipette puller and filled by boiling in 3 M KCl under reduced atmospheric pressure. The tip resistance of the electrodes was between 10 and 30 Megohms. The microelectrode were connected through an Ag/AgCl bridge to Bioelectric Instruments VHI holders and to the input of a unity gain voltage follower with an input impedance of $>10^{11}$ ohms. The outputs from these preamplifiers were displayed on a Tektronix 564 storage oscilloscope. Photographs were taken with a C 12 Polaroid camera. The preparation was stimulated through a bipolar extracellular electrode insulated with Teflon except at its tip. Stimulating pulses were derived

from a Grass Model S 4 pulse generator and isolation unit. The pulse generator was triggered by a series of Tektronix 161 pulse generators. This system provided pulses at any desired basic driving frequency and a dividing circuit between two pulse generators permitted the delivery of premature test pulses after every sixth basic pulse at any desired coupling interval. The pulses were 3 msec in duration and adjusted to two times threshold.

In each experiment two simultaneous action potentials were recorded. Initially while pacing the preparation from the proximal right bundle branch one microelectrode was moved in 1 to 2 mm steps from the proximal right bundle out along a false tendon and distal Purkinje fibers up to the point where they penetrated the endocardial surface of the right ventricular free wall. At each position action potential duration was recorded. After determining the area of maximal action potential duration one microelectrode was positioned approximately 1 cm proximal to that area and a second microelectrode was placed 2 to 3 mm distal to the area. Propagation of premature test pulses was then examined. Finally, normal Tyrode solution was substituted with Tyrode solution containing lidocaine hydrochloride in concentrations from 1×10^{-8} to $1 \times 10^{-4} \text{M}$.

Definitions

1 *Action potential duration (APD)* The interval between the upstroke of the action potential (phase 0) and 80 per cent repolarization.

2 *Gating point* The site in the false tendon at which maximal action potential duration was observed.

3 *Functional refractory period of the gate (FRPG)* The minimal interval between the upstrokes of action potentials distal to the gating point in response to a premature stimulus delivered proximal to the gate.

Results

The effect of lidocaine on the time course of repolarization in Purkinje fibers was examined in 10 preparations. The results of these studies are presented in Fig. 2. In each study the preparation was driven at a frequency of 60 beats per minute (i.e. cycle length 1000 msec) and transmem-

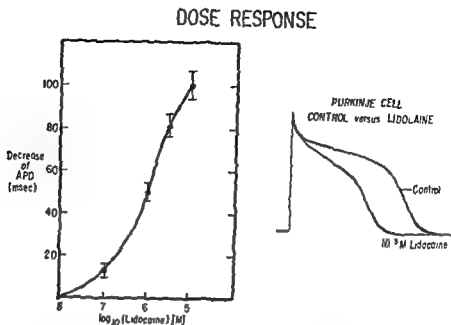


Fig 2 Effects of lidocaine on action potential (right panel) and dose-response curve (left panel) APD = action potential duration

brane action potentials were recorded from Purkinje fibers in the free running strand approximately 1 cm from its insertion into the subendocardium of the right ventricle. Lidocaine produced a dose dependent decrease of action potential duration. A typical transmembrane action potential obtained before and after exposure to lidocaine is shown in the right panel of Fig 2. The relation between the concentration of lidocaine and the decrease of action potential duration is shown in the left panel of Fig 2. At $1 \times 10^{-8} \text{ M}$ lidocaine produced no change of action potential duration. At $1 \times 10^{-7} \text{ M}$ lidocaine decreased action potential duration by $13.3 \pm 1.5 \text{ msec}$. At 1×10^{-6} and $5 \times 10^{-6} \text{ M}$ lidocaine decreased action potential duration by 50 ± 4.5 and $81 \pm 5.0 \text{ msec}$ respectively. At a concentration of $1 \times 10^{-5} \text{ M}$ lidocaine decreased action potential duration by $100 \pm 6.7 \text{ msec}$ and no further decrease was noted at $5 \times 10^{-5} \text{ M}$. Previous work by Bigger and Hessebutter⁴ and by Gianelly and colleagues⁵ has indicated that the therapeutic concentration of lidocaine in plasma is approximately $2.5 \mu\text{g}$ per milliliter. Lidocaine $1 \times 10^{-5} \text{ M}$ corresponds to a concentration of $2.34 \mu\text{g}$ per milliliter.

Under control conditions the duration of action potentials recorded from Purkinje fibers along the right bundle branch varies. As cells were impaled sequentially in the

proximal right bundle branch, false tendon, and terminations of the Purkinje system at the endocardial surface of the right ventricle, action potential duration increased and then decreased. The region of maximal action potential duration was located within the false tendon and was usually 2 to 5 mm proximal to the site where a free running strand penetrated the endocardial surface. A representative map of action potential durations along the right bundle branch and false tendon before and after exposure to lidocaine is shown in Fig 3. Under control conditions action potential duration increased from 225 msec in the proximal right bundle branch to 240 msec at the level of the anterior papillary muscle. As the right bundle branch became a free running strand, action potential duration increased to a maximum of 305 msec and then decreased rapidly to 200 msec just before the strand penetrated the endocardium of the right ventricular free wall. The preparation was then perfused with Tyrode solution which contained $1.5 \times 10^{-5} \text{ M}$ lidocaine. The steady state maximal effects of lidocaine were achieved in 7 to 8 minutes. Twenty minutes after exposure to lidocaine the map of action potential duration over the conducting system was repeated. Action potential duration was decreased by lidocaine throughout the preparation, but the maximal effect was

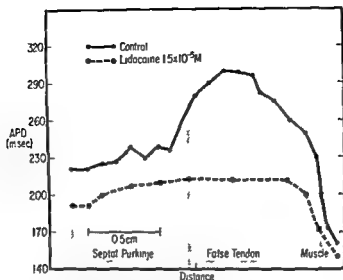


Fig. 3 Map of action potential durations in the right bundle branch, false tendon, and ventricular muscle (see Fig. 1). Solid circles show APD before perfusion with lidocaine. Dashed curve shows APD after exposure to lidocaine. Note that lidocaine shortened APD maximally in the false tendon.

Table 1 Effects of lidocaine ($1 \times 10^{-3} M$) on action potential duration (msec)

| Experiment no | Proximal RBB | | Mid RBB | | Gate | | Distal | | Ventricular muscle | |
|----------------|--------------|-------|------------|-------|-------------|-------|------------|-------|--------------------|-------|
| | Before | After | Before | After | Before | After | Before | After | Before | After |
| 1 | 275 | 220 | 290 | 225 | 300 | 220 | 235 | 200 | 180 | 175 |
| 2 | 205 | 200 | 265 | 210 | 330 | 210 | 250 | 210 | 150 | 145 |
| 3 | 210 | 215 | 260 | 220 | 320 | 220 | 230 | 198 | 160 | 155 |
| 4 | 270 | 210 | 260 | 227 | 295 | 235 | 240 | 220 | 143 | 134 |
| 5 | 235 | 215 | 265 | 270 | 310 | 220 | 240 | 200 | 155 | 150 |
| 6 | 275 | 200 | 240 | 210 | 305 | 210 | 235 | 200 | 170 | 160 |
| 7 | 215 | 195 | 240 | 195 | 370 | 195 | 260 | 190 | 180 | 175 |
| 8 | 210 | 200 | 260 | 210 | 305 | 210 | 240 | 210 | 160 | 150 |
| 9 | 220 | 205 | 240 | 205 | 330 | 200 | 250 | 200 | 180 | 195 |
| 10 | 230 | 200 | 250 | 205 | 305 | 200 | 250 | 200 | 175 | 168 |
| Average change | 13.5 = 3.3 | | 47.8 = 4.4 | | 100.0 = 6.7 | | 40.2 = 4.4 | | 6.6 = 0.8 | |
| P value | P > 0.001 | | P > 0.001 | | P > 0.001 | | P > 0.001 | | P > 0.001 | |

noted at the region of maximal action potential duration. Thus in the proximal right bundle branch, action potential duration was decreased from 225 to 200 msec while in the free running strand action potential duration decreased from 290 to 205 msec. In the most distal Purkinje fibers, where action potential duration was already short, the effects of lidocaine were small relative to the free running strand. The effects of lidocaine on the duration of action potentials in 10 preparations are

summarized in Table I. Lidocaine decreased action potential duration in the proximal right bundle branch by an average of 13 msec at the base of the papillary muscle by 47 msec at the area of maximum action potential duration by 100 msec and at distal fibers by 40 msec. Lidocaine decreased action potential duration of ventricular muscle by only 6 msec. Thus the most prominent effect of lidocaine was to shorten action potential duration at the region of maximal duration during the control period.

DOSE RESPONSE

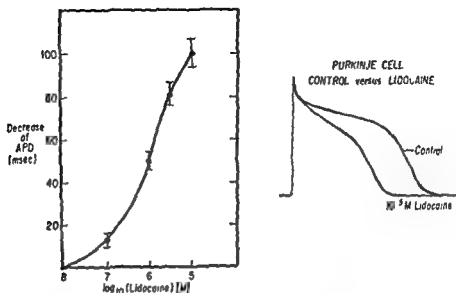


Fig 2 Effects of lidocaine on action potential (right panel) and dose response curve (left panel) APD = action potential duration

brane action potentials were recorded from Purkinje fibers in the free running strand approximately 1 cm from its insertion into the subendocardium of the right ventricle. Lidocaine produced a dose dependent decrease of action potential duration. A typical transmembrane action potential obtained before and after exposure to lidocaine is shown in the right panel of Fig 2. The relation between the concentration of lidocaine and the decrease of action potential duration is shown in the left panel of Fig 2. At 1×10^{-8} M, lidocaine produced no change of action potential duration. At 1×10^{-7} M lidocaine decreased action potential duration by 13.3 ± 1.5 msec. At 1×10^{-6} and 5×10^{-6} M lidocaine decreased action potential duration by 50 ± 4.5 and 81 ± 5.0 msec, respectively. At a concentration of 1×10^{-5} M lidocaine decreased action potential duration by 100 ± 6.7 msec and no further decrease was noted at 5×10^{-5} M. Previous work by Bigger and Hennessy⁴ and by Ganelly and colleagues⁵ has indicated that the therapeutic concentration of lidocaine in plasma is approximately $2.5 \mu\text{g}$ per milliliter. Lidocaine 1×10^{-5} M corresponds to a concentration of $2.34 \mu\text{g}$ per milliliter.

Under control conditions the duration of action potentials recorded from Purkinje fibers along the right bundle branch varies. As cells were impaled sequentially in the

proximal right bundle branch, false tendon, and terminations of the Purkinje system at the endocardial surface of the right ventricle, action potential duration increased and then decreased. The region of maximal action potential duration was located within the false tendon and was usually 2 to 5 mm proximal to the site where a free running strand penetrated the endocardial surface. A representative map of action potential durations along the right bundle branch and false tendon before and after exposure to lidocaine is shown in Fig 3. Under control conditions action potential duration increased from 220 msec in the proximal right bundle branch to 240 msec at the level of the anterior papillary muscle. As the right bundle branch became a free running strand action potential duration increased to a maximum of 305 msec and then decreased rapidly to 200 msec just before the strand penetrated the endocardium of the right ventricular free wall. The preparation was then perfused with Tyrode solution which contained 1.5×10^{-5} M lidocaine. The steady state maximal effects of lidocaine were achieved in 7 to 8 minutes. Twenty minutes after exposure to lidocaine the map of action potential duration over the conducting system was repeated. Action potential duration was decreased by lidocaine throughout the preparation but the maximal effect was

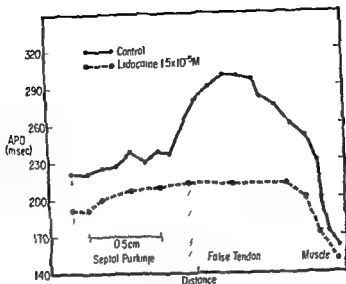


Fig. 3 Map of action potential durations in the right bundle branch, false tendon, and ventricular muscle (see Fig. 1) solid circles show APD before perfusion with lidocaine. Dashed curve shows APD after exposure to lidocaine. Note that lidocaine shortened APD maximally in the false tendon.

Table I Effects of lidocaine ($1 \times 10^{-5} M$) on action potential duration (msec)

| Experiment no | Proximal RBB | | Mid RBB | | Gate | | Distal | | Ventricular muscle | |
|----------------|----------------|-------|----------------|-------|-----------------|-------|----------------|-------|--------------------|-------|
| | Before | After | Before | After | Before | After | Before | After | Before | After |
| 1 | 223 | 220 | 290 | 225 | 300 | 220 | 235 | 200 | 180 | 175 |
| 2 | 205 | 200 | 265 | 210 | 330 | 210 | 250 | 210 | 150 | 145 |
| 3 | 210 | 215 | 260 | 220 | 320 | 220 | 230 | 198 | 160 | 155 |
| 4 | 220 | 210 | 260 | 222 | 295 | 235 | 240 | 220 | 143 | 134 |
| 5 | 235 | 215 | 265 | 220 | 310 | 220 | 240 | 200 | 155 | 150 |
| 6 | 225 | 200 | 240 | 210 | 405 | 210 | 235 | 200 | 170 | 160 |
| 7 | 215 | 195 | 270 | 195 | 320 | 195 | 260 | 190 | 180 | 175 |
| 8 | 210 | 200 | 260 | 210 | 405 | 210 | 240 | 210 | 160 | 150 |
| 9 | 220 | 205 | 240 | 205 | 330 | 200 | 250 | 200 | 180 | 195 |
| 10 | 230 | 200 | 250 | 205 | 405 | 200 | 250 | 200 | 175 | 165 |
| Average change | 13.5 \pm 3.3 | | 47.8 \pm 4.4 | | 100.0 \pm 6.7 | | 40.2 \pm 4.4 | | 6.6 \pm 0.8 | |
| P value | P > 0.001 | | P > 0.001 | | P > 0.001 | | P > 0.001 | | P > 0.001 | |

noted at the region of maximal action potential duration. Thus in the proximal right bundle branch action potential duration was decreased from 225 to 200 msec while in the free running strand action potential duration decreased from 290 to 205 msec. In the most distal Purkinje fibers where action potential duration was already short the effects of lidocaine were small relative to the free running strand. The effects of lidocaine on the duration of action potentials in 10 preparations are

summarized in Table I. Lidocaine decreased action potential duration in the proximal right bundle branch by an average of 13 msec at the base of the papillary muscle by 47 msec at the area of maximum action potential duration by 100 msec and at distal fibers by 40 msec. Lidocaine decreased action potential duration of ventricular muscle by only 6 msec. Thus the most prominent effect of lidocaine was to shorten action potential duration at the region of maximal duration during the control period.

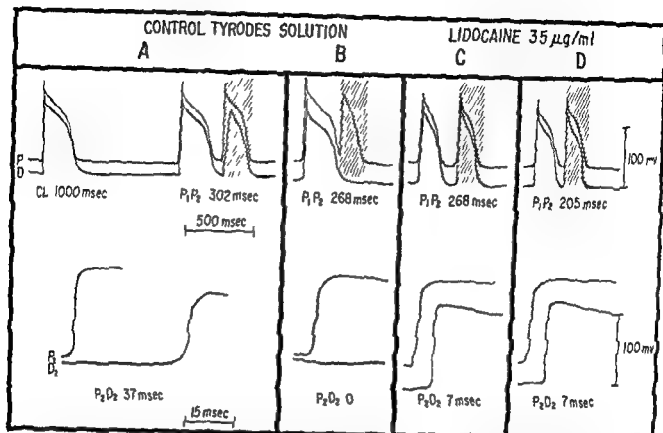


Fig 4 A through D Effects of lidocaine on APD and propagation of early premature beats P = cell proximal to the gate D = cell distal to the gate Lower tracings show expanded time sweep to illustrate dV/dt of the upstroke and P_1D_1 interval Panels A and B are before lidocaine Panels C and D are after exposure to lidocaine See text for details

As a consequence of this effect action potential durations became more uniform in the Purkinje strand and the marked discrepancy of action potential durations which normally exists between the free running strand and the adjacent muscle was largely abolished.

Myerburg and colleagues¹² described the gradual increase of action potential duration in distal Purkinje fibers and noted that the area of maximal duration functioned as a "gate." Premature stimuli delivered either to the proximal right bundle branch or to distal ventricular muscle were blocked at the gate when the coupling interval was less than action potential duration at the gate. Fig 4 (panels A and B) illustrates this phenomenon. In panel A action potentials were recorded simultaneously from cells proximal and distal to the area of maximal action potential duration. After a series of beats at a basic cycle length of 1 000 msec a premature stimulus was introduced into the right bundle branch. In the proximal cell the premature response occurred 302

msec after the last basic response and propagated across the gate to the distal cell. The interval between the proximal and distal response on the premature beat was 37 msec. At a coupling interval of 301 msec the premature pulse elicited a response in the proximal cell, but this response failed to propagate across the gate. A premature test pulse still elicited a response in the proximal cell at coupling intervals as short as 268 msec (panel B). The effects of lidocaine are shown in panels C and D. After lidocaine action potential duration was reduced. At a coupling interval of 268 msec (the same as panel B) a premature test pulse elicited a response in the proximal cell which propagated across the gate. The interval between the proximal and distal responses on the premature beat was only 7 msec (panel C). After exposure to lidocaine the coupling interval between the basic and premature beat could be further reduced to 205 msec while propagation across the gate was maintained (panel D). Not only was propagation across the gate

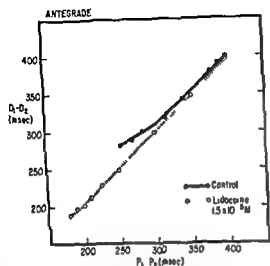


Fig 5 Effects of lidocaine on the functional refractory period of the gate (antegrade) $P_1 P_2$ = interval between basic and premature responses proximal to gate $D_1 D_2$ = corresponding intervals distal to the gate Antegrade = propagation from right bundle to right ventricle Basic cycle length = 1 000 msec

maintained but the decremental conduction and delay which was evident in panel A was absent after lidocaine (panels C and D) Results comparable to these were observed in each of 10 preparations

The effects of lidocaine on propagation across the gate are further depicted in Figs 5 and 6 In Fig 5 action potentials were recorded proximal and distal to the gate while driving the preparation from the right bundle branch (antegrade) Premature test pulses were delivered to the right bundle at progressively shorter coupling intervals and the interval between action potentials in the proximal cell ($P_1 P_2$) was plotted against the corresponding interval between action potentials in the distal cell ($D_1 D_2$) During the control period the $D_1 D_2$ interval began to increase at $P_1 P_2$ intervals of approximately 300 msec (i.e. delayed propagation of the premature response) At a $P_1 P_2$ interval of 250 msec the premature beat failed to propagate across the gate The shortest interval between responses in the distal cell (i.e. 270 msec in Fig 5) defined the functional refractory period of the gate After lidocaine the functional refractory period of the gate was reduced to 190 msec Decremental conduction of early premature

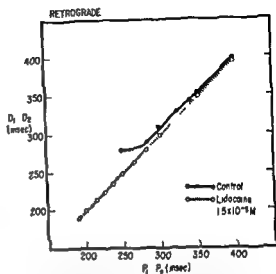


Fig 6 Effects of lidocaine on the functional refractory period of the gate (retrograde) Nomenclature the same as Fig 5 Retrograde = propagation from right ventricle to right bundle branch Basic cycle length = 1 000 msec

beats which was evident before lidocaine was abolished

In Fig 6 the same type of plot was used to depict the effects of lidocaine on retrograde conduction across the gate In this instance the preparation was driven from the right ventricular free wall so that the impulse propagated retrograde across the free running strand to the right bundle branch Action potentials were recorded proximal (near ventricular muscle) and distal (near the right bundle branch) to the area of maximal action potential duration Before lidocaine the functional refractory period of the gate was 270 msec and premature beats with short coupling intervals were propagated with decremental conduction After lidocaine the functional refractory period was reduced to 190 msec and decremental conduction was abolished

Discussion

In 1969 our group reported certain of the electrophysiologic effects of lidocaine on the heart in awake dogs¹¹ Lidocaine decreased the ventricular rate of dogs with chronic heart block and produced a substantial increase in the threshold for stimulation of atrial and ventricular muscle Somewhat to our surprise lidocaine failed

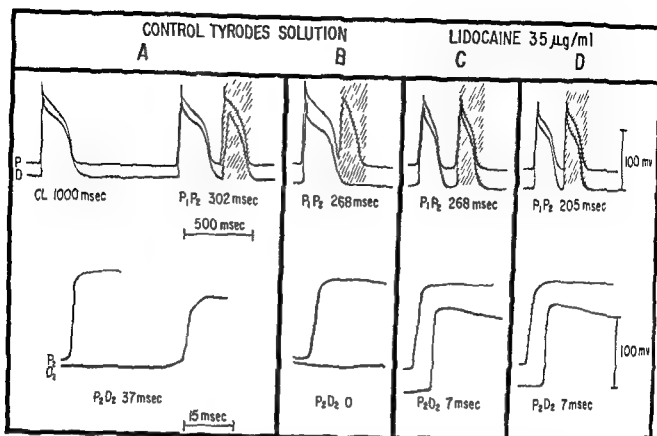


Fig. 4 A through D Effects of lidocaine on APD and propagation of early premature beats. P = cell proximal to the gate. D = cell distal to the gate. Lower tracings show expanded time sweep to illustrate dv/dt of the upstroke and P_2D_2 interval. Panels A and B are before lidocaine. Panels C and D are after exposure to lidocaine. See text for details.

As a consequence of this effect action potential durations became more uniform in the Purkinje strand and the marked discrepancy of action potential durations which normally exists between the free running strand and the adjacent muscle was largely abolished.

Myerburg and colleagues¹³ described the gradual increase of action potential duration in distal Purkinje fibers and noted that the area of maximal duration functioned as a 'gate'. Premature stimuli delivered either to the proximal right bundle branch or to distal ventricular muscle were blocked at the gate when the coupling interval was less than action potential duration at the gate. Fig. 4 (panels A and B) illustrates this phenomenon. In panel A action potentials were recorded simultaneously from cells proximal and distal to the area of maximal action potential duration. After a series of beats at a basic cycle length of 1 000 msec a premature stimulus was introduced into the right bundle branch. In the proximal cell the premature response occurred 302

msec after the last basic response and propagated across the gate to the distal cell. The interval between the proximal and distal response on the premature beat was 37 msec. At a coupling interval of 301 msec the premature pulse elicited a response in the proximal cell but this response failed to propagate across the gate. A premature test pulse still elicited a response in the proximal cell at coupling intervals as short as 268 msec (panel B). The effects of lidocaine are shown in panels C and D. After lidocaine action potential duration was reduced. At a coupling interval of 268 msec (the same as panel B) a premature test pulse elicited a response in the proximal cell which propagated across the gate. The interval between the proximal and distal responses on the premature beat was only 7 msec (panel C). After exposure to lidocaine the coupling interval between the basic and premature beat could be further reduced to 205 msec while propagation across the gate was maintained (panel D). Not only was propagation across the gate

maximal in distal Purkinje fibers at the region of the gate. The effect of lidocaine on proximal Purkinje fibers within the right bundle branch and on ventricular muscle is much less marked. Thus one consequence of the action of lidocaine is to diminish nonuniform repolarization in distal Purkinje fibers and to reduce the discrepancy of action potential durations which normally exists between Purkinje fibers and ventricular muscle. A second consequence of this action is to abolish or markedly attenuate the gating function of the distal Purkinje fibers. An important result of these actions is to shorten the functional refractory period of the gate and to abolish decremental conduction of premature impulses which arrive at the gating segment early in its phase of recovery of excitability. Our studies indicate that these actions of lidocaine on the gating segment of Purkinje tissue are applicable whether the premature impulse invades the gate from above (i.e. antegrade) or from below (i.e. retrograde).

In considering the potential significance of our data to the intact heart at least three pieces of additional data would be desirable. First it remains to be proved that lidocaine exerts the above actions on the Purkinje system of the intact awake animal or man. Second it would be very desirable to know if lidocaine exerts a quantitatively similar effect on all gating segments since as noted by Myerburg and colleagues¹¹ nonuniformity of gates whether spontaneous or pharmacologically induced could enhance the likelihood of arrhythmias rather than the protective effect we have attributed to the actions of lidocaine. Finally it would be of particular interest to induce re-entrant beating at the Purkinje muscle junction by interventions which are analogous to those which may operate in disease and to examine the influence of lidocaine under these conditions.

For the reasons noted above it is premature to ascribe the antiarrhythmic actions of lidocaine to its effect on distal Purkinje fibers and the gating function of that segment of the Purkinje muscle junction. The marked discrepancies of action potential duration which normally exist in this region however as well as the proximity of this segment to areas of disease—

ie intramural infarcts—and the apparent sensitivity of the gating segment to pharmacological agents all make this a particularly attractive model system for the evaluation of antiarrhythmic drugs.

Summary

These studies were designed to examine the electrophysiological effects of lidocaine on distal Purkinje fibers of the canine heart. Standard microelectrode techniques were used to record transmembrane action potentials from the right bundle branch free running strands of Purkinje tissue, and ventricular muscle. The duration of action potentials was maximal in the distal Purkinje fibers. The long duration of action potentials in these distal fibers determined the functional refractory period of the Purkinje muscle junction. Early premature beats initiated on either side of the distal Purkinje fibers propagated with decrement or were blocked. Lidocaine shortened the action potential of Purkinje fibers but had little effect on muscle. The effect of lidocaine on action potential duration was dose dependent and was maximal in these distal Purkinje fibers. As a consequence of its ability to shorten action potential duration lidocaine reduced the degree of nonuniformity of recovery of excitability. Lidocaine shortened the functional refractory period of distal Purkinje fibers and abolished decremental conduction of early premature beats. These effects may contribute to the antiarrhythmic actions of lidocaine.

REFERENCES

- Southworth J L, McKusick V A, Pierce E C and Rawson F L. Ventricular fibrillation. *JAMA* 143:717 1950.
- Lown B, Fakhro A M, Hood W B and Thorn G W. The coronary care unit: New perspectives and directions. *JAMA* 199:188 1967.
- Chopra M P, Portal R W and Aber C P. Lignocaine therapy after acute myocardial infarction. *Br Med J* 1:713 1969.
- Bigger J T Jr and Heissenbuttel R H. The use of procaine amide and lidocaine in the treatment of cardiac arrhythmias. *Progr Cardiovasc Dis* 11:515 1969.
- Giannelis J I, Von Der Groben J O, Spivack A I and Harrison D C. Effect of lidocaine on ventricular arrhythmias in patients with coronary heart disease. *N Engl J Med* 277:1215 1967.
- Bedell G N, Culbertson J W and Ehren

to produce significant alterations of conduction velocity in Purkinje tissue total ventricular activation time or refractoriness of ventricular muscle. Lidocaine thus appeared to have very different electrophysiological properties than quinidine.¹⁴ At the time of our earlier report there were no published data concerning the effects of lidocaine on the action potential of cardiac fibers.

In 1969 Davis and Temte¹ described the effects of lidocaine on isolated Purkinje fibers and ventricular muscle. At concentrations from 5 to 50 μg per milliliter the most prominent effect of lidocaine was an acceleration of repolarization which resulted from an increase in the slope of phase 2 and an earlier onset of phase 3. This effect of lidocaine was confined to the Purkinje fibers; however, and little change in the duration of action potentials was noted in ventricular muscle. These investigators¹ also observed that lidocaine abolished decremental conduction across the distal Purkinje fibers when premature stimuli were delivered to the preparation during the early phase of recovery of excitability.

In 1970 Bigger and Mandel¹⁵ reported that lidocaine in concentrations of 1×10^{-6} and 1×10^{-5} attenuated or abolished spontaneous phase 4 diastolic depolarization in Purkinje fibers and shortened action potential duration. Lidocaine had little or no effect on the maximal rate of phase 0 depolarization. Bigger and Mandel¹⁵ postulated that the effect of lidocaine on phase 4 diastolic depolarization and on the duration of the action potential could be attributed to an increase of membrane conductance for potassium.

In 1971, Singh and Vaughan Williams¹⁶ confirmed the observations of Davis and Temte¹ and Bigger and Mandel¹⁵ with respect to the effect of lidocaine on action potential duration. They also indicated that lidocaine decreased $\text{max } dv/dt$ of the upstroke of cardiac muscle. They considered that the failure of Bigger and Mandel¹⁵ to observe this latter effect might have been accounted for by a relatively low K^+ concentration in their Tyrode solution. Most investigators now agree, however, that an important and consistent effect of lidocaine is to decrease action potential duration

that this effect is more prominent in Purkinje fibers than in ventricular muscle and that at therapeutic concentrations lidocaine has only a slight depressant effect on conduction velocity.

Moe and Han and their colleagues¹⁷⁻¹⁹ have stressed the potential importance of nonuniform recovery of excitability in adjacent regions of myocardium to the genesis of reentrant arrhythmias. In the above reports these investigators have shown that the 'vulnerable period' corresponds to an interval of nonhomogenous recovery of excitability in the myocardium and that agents which either increase or decrease the degree of nonhomogenous recovery respectively make the heart more or less vulnerable to fibrillation induced by a premature stimulus. The data which led to that conclusion were based on measurements of the refractory period at adjacent areas of epicardial muscle, and discrepancies in the range of 10 to 20 msec. In the report by Myerburg and colleagues²⁰ the duration of the action potential and refractory period of distal Purkinje fibers was noted to be 80 to 100 msec longer than in adjacent muscle. Myerburg and colleagues²¹ attributed a protective effect to the gating segment of distal Purkinje fibers, but considered the possibility that should a localized breakdown of the gate occur during ischemia for example that this could create certain of the conditions necessary to support a reentry circuit. Previous experiments by Alnis and co-workers²² had demonstrated slow conduction and unidirectional block at the Purkinje muscle junction. The observations by Moe and colleagues¹⁷⁻¹⁹, Myerburg and associates²¹ and Alnis and co-workers²² suggested to us that the marked differences of action potential duration at the Purkinje muscle junction might contribute to arrhythmias of the vulnerable period and that distal Purkinje fibers might be a particularly rewarding site to examine the effects of lidocaine.

Our studies have shown that lidocaine in concentrations which are equivalent to therapeutic levels in plasma of man exerts a prominent effect on the duration of action potentials in Purkinje fibers. The ability of lidocaine to shorten action potential duration is dose dependent and the effect is

Activation of the human fetal heart

Antonio Brusca MD

Erennio Rosettani MD

Turin Italy

Recent attempts to correlate electrical activity of the heart to the surface electrical field^{1,2} and the new data on disturbances of impulse transmission in the ventricles³ have aroused a new interest in cardiac activation.

Our present knowledge of the excitation process of the heart is based mainly on animal experiments.⁴⁻⁶ Some data also have been gathered in humans⁷⁻¹⁰ mostly from studies performed during cardiac surgery.¹¹⁻¹³ These studies, although useful, have obvious technical limitations. The work of Durrer and his group¹⁴ on the perfused human heart is an important exception; these workers have also demonstrated that, as far as cardiac excitation is concerned, the electrical behavior of the isolated and perfused heart is practically identical to the one of the heart *in vivo*.

This paper deals with a study of the activation process of the human fetal heart.

Material and methods

The present study was carried out on 11 human fetal hearts removed within 30 minutes after death, which were revived and perfused with the Langendorf technique. The age of the fetuses varied from 5 1/2 to 9 months. In each case a routine ECG was taken shortly before death and was found to be normal.¹⁵ At the end of

each experiment the heart was carefully examined and was found to be free from any gross structural abnormality. Perfusion was performed with a modified Tyrode solution.*

By means of surface and multipolar needle electrodes with an interterminal distance of 0.5 mm, bipolar records were obtained from the epicardium, left and right endocardium, and from the septal wall. An eight channel Electronics for Medicine apparatus was used with an electronic switch at a paper speed of 200 mm per second. Magnification of the tracings with magnifying lens allowed measurement of activation times with an accuracy of 1 msec. An average of 300 points were explored in each heart (of these between 50 and 60 were on the atria). A reference lead was recorded with one electrode placed at the apex and the other at the root of the aorta. The left endocardium was exposed through a longitudinal cut along the obtuse margin of the heart. Similarly, the right ventricular cavity was opened with a cut along the acute margin. The results were discarded if any changes in the form and duration of the QRS occurred in the reference lead, which was carefully monitored during the experimental procedure. For atrial activation, the beginning of the P wave served as a time reference for ven-

* From the Di. Scienze di Cardiologia, Clinica Medica Università di Torino, Torino, Italy.

Received for publication Aug 15 1972.

Reprints requested to: A. Brusca, MD, Via Meravigli 11, Torino, Italy.

Tyrode solution = NaCl 137 mm, KCl 2.7 mm, CaCl₂ 2.7 mm, NaHCO₃ 12 mm, NaH₂PO₄ 1.8 mm, MgCl₂ 0.3 mm, and glucose 5.5 mm.

- haft J L Management of patients with mitral stenosis before during and after mitral valvuloplasty *Arch Intern Med* 94:718 1954
- 7 Likoff W Cardiac arrhythmias complicating surgery *Am J Cardiol* 3:427 1959
- 8 Weiss A Intravenous use of lidocaine for ventricular arrhythmias *Anesth Analg (Cleve)* 39:369 1960
- 9 Morrow D H and Bosomworth P P Anesthesia and digitalis toxicity III Effect of lidocaine on ouabain induced ventricular tachycardia *Anesth Analg (Cleve)* 46:183 1967
- 10 Grossman J I Lubow I A Frieden J and Rubin I L Lidocaine in cardiac arrhythmias *Arch Intern Med* 121:396 1968
- 11 Sugimoto T Schaal S F Dunn N M and Wallace A G Electrophysiological effects of lidocaine in awake dogs, *J Pharmacol Exp Ther* 166 146 1969
- 12 Davis L D and Temte J V Electrophysiological actions of lidocaine on canine ventricular muscle and Purkinje fibers *Circ Res* 24:639 1969
- 13 Myerburg R J Stewart J W and Hoffman B F Electrophysiological effects of canine peripheral AV conducting system *Circ Res* 26:361 1970
- 14 Wallace A G Cline R E Sealy W C Young W G and Troyer W G Electrophysiological effects of Quinidine Studies using chronically implanted electrodes in awake dogs with and without cardiac denervation *Circ Res* 19:960 1966
- 15 Bigger J T Jr and Mandel W J Effect of lidocaine on electrophysiological properties of ventricular muscle and Purkinje fibers *J Clin Invest* 49:63, 1970
- 16 Singh B N and Vaughan Williams E M Effect of altering potassium concentration on the action of lidocaine and diphenylhydantoin on rabbit atrial and ventricular muscle *Circ Res* 29 786 1971
- 17 Moe G K On the multiple wavelet hypothesis of atrial fibrillation *Arch Int Pharmacodyn* 140:183 1962
- 18 Han J and Moe G K Nonuniform recovery of excitability in ventricular muscle *Circ Res* 14:44 1964
- 19 Han J DeJalon P G and Moe G K Adrenergic effects on ventricular vulnerability *Circ Res* 14 516 1964
- 20 Alanis J and Benitez D Transitional potentials and the propagation of impulses through different cardiac cells in Sano T Mizuhira V and Matsuda K editors *Electrophysiology and ultrastructure of the heart Tokyo Japan 1967 Bunlendo Co Ltd*

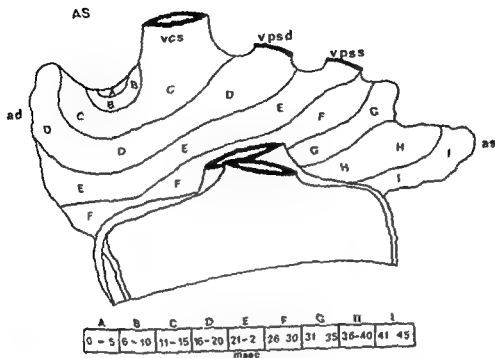


Fig 2 Same heart as in Fig 1 AS = anterior superior a pect of the atria ad = right atrium as = left atrium vcs = superior vena cava vpsd = right superior pulmonary vein vpsl = left superior pulmonary vein

several exceptions. In some instances the latest activity occurs on the posterior lateral portion of the left ventricle while excitation of the subpulmonic area occurs relatively early. Furthermore in the case presented in Fig 3C a small area of very early activation is present on the posterior aspect of the right ventricle and another area of early activity is observed on the upper anterior portion of the left ventricle near the AV groove.

Altogether epicardial activation when a sufficient number of points are explored seems to be a phenomenon which is highly complex and extremely variable from one normal heart to the other.

This observation cast some doubts on the reliability of predicting the details of epicardial and ventricular excitation from mapping of surface potential differences. Similar observations have been previously reported by King and colleagues²⁵ for atrial activation.

Within the ventricles three regions of very early activity are found (Fig 4A) all of them on the left endocardial surface. The first one is located on the central por-

tion of the middle third of the septum the second area is high on the parietal endocardium close to the anterior AV groove and the third one again on the parietal endocardium is located near the mid lower portion of the left posterior paraseptal region. These findings are in complete agreement with those obtained by Durrer and associates¹ in adult human hearts. The three areas mentioned above might well represent the functional trifurcation of the left main bundle.

From these initial foci activity spreads quickly in the next 5 msec to most of the left parietal and septal endocardium. The anterior apical and basal septal zones are the last activated regions. Excitation on the left endocardium seems to be a rather uniform and quick event which is completed within 20 msec from the beginning of the QRS complex.

In the right ventricular endocardium (Fig 4B) activity occurs first in just one area on the parietal wall near the base of the anterior right papillary muscle 15 msec after the beginning of QRS. From this region activity spreads slowly along

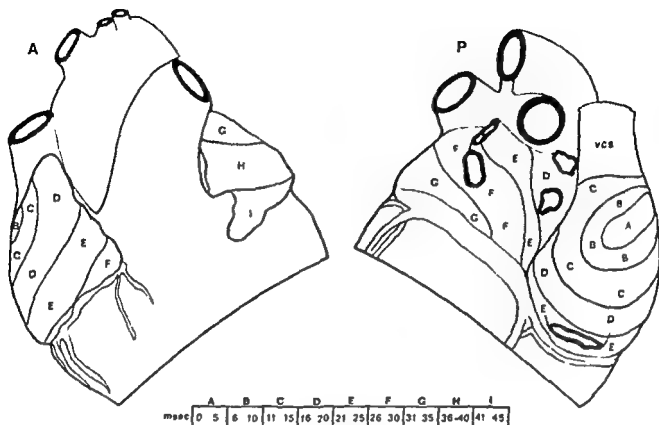


Fig 1 Activation of the atria (Age of subject 61 $\frac{1}{2}$ months weight of heart 4.8 grams) A = anterior view P = posterior view VCS = superior vena cava Activation times are given in msec after the beginning of the P wave in the reference lead

tricular activation the beginning of QRS was taken as time zero

Results and discussion

Activation of the atria begins in the region of the sinus node as an irregular oval area, within 0 and 5 msec from the beginning of the P wave in the control lead. From this area excitation spreads radially following a series of concentric equal isochrones. Each isochrone involves simultaneously both a posterior and a superior anterior portion of the atrial wall. As shown in Figs 1 and 2 the activation front proceeds in a cranio caudal direction and from right to left. In four cases no electrical activity could be registered on the posterior aspect of the left atrium in the region of the pulmonary veins. The width of the isochrones is equal in each region of the atria suggesting a rather uniform velocity of conduction. This finding and the general pattern of atrial activation would tend to exclude the existence of specialized atrial pathways along which activity spreads preferentially

from right to left atrium. Consequently it should be pointed out that the idea of functional specialized interatrial tracts is based mainly on indirect deductions in animal experiments³ by the difference in shape of intracellular action potentials²⁶ and from the sensitivity of atrial cells to changes in external potassium concentration.²⁷ However, since the atrial septum was not explored in our studies the existence of possible internodal preferential tracts can not be ruled out.

Epicardial activation of the ventricles begins as a rule in a small area on the trabecular wall of the right ventricle (Fig 3A) it then proceeds to the anterior and posterior paraseptal regions. Sometimes as in the case shown in Fig 3B this second area of early activation involves the apical region as well. Epicardial breakthrough subsequently occurs in an apico basal direction involving large portions of the free ventricular wall. The first activated regions are the diaphragmatic and basal portions of the right ventricle. However there are

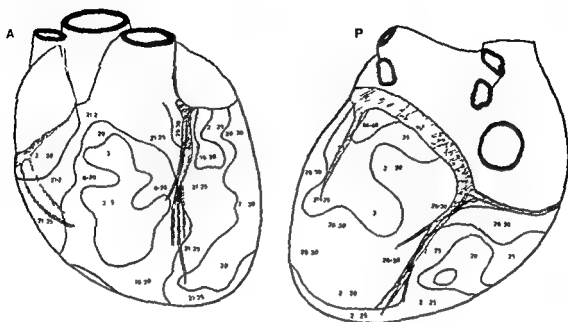


Fig 1C Epicardial activation of the ventricles of a fetal human heart (Age of subject 7 months heart weight 6 grams) Abbreviations as in 1A Note the early activity on the posterior aspect of the right ventricle and on the anterior part of the left ventricle near the AV sulcus

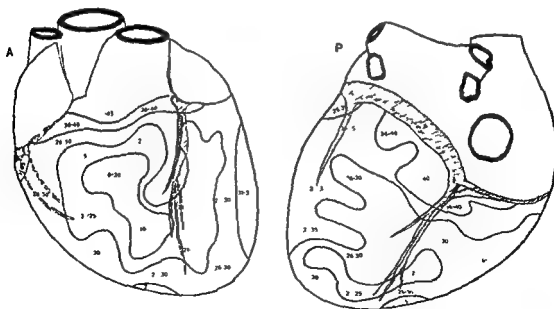


Fig 1D Epicardial activation of the ventricles of a fetal human heart (Age of subject 6 1/2 months heart weight 5.5 grams) Abbreviations as in 1A

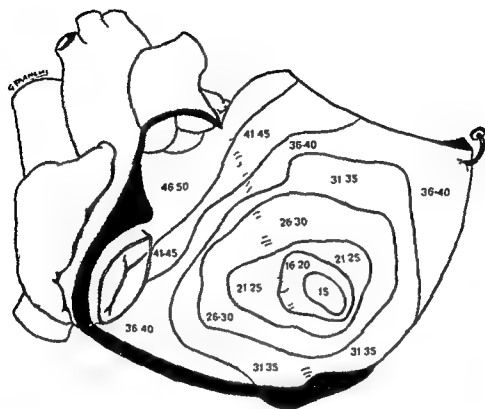


Fig 4B Same heart as illustrated in 4A Activation of the right endocardium

of the septum while the wave of excitation advances mainly tangentially in the apical and basal septal mass. Excitation of the septum always occurred from left to right right to left penetration as described in dogs²⁹ and in adult human hearts³⁰ was never observed.

Summary

Ipicardial activation of atria and ventricles endocardial activation of the ventricles and intramural activation of the interventricular septum were studied in 11 isolated, revived and perfused human fetal hearts. No evidence was found for specialized pathways in the atria. Earliest epicardial breakthrough in the ventricles occurred in the area pretrabecularis of the right ventricle. Sometimes an early area was found on the posterior aspect of the right ventricle. Latest activated areas were the diaphragmatic and basal aspects of the right ventricle. On occasion the latest

region was the posterolateral part of the left ventricle. On the endocardial surface three regions of early activity were found: the central part of the middle third of the left part of the septum, an area located near the base of the left anterior free wall and a region near the middle lower part of the left posterior paraseptal region. Septal activation occurred in a left to right direction only; nowhere was a right to left spread of excitation found. The intramural spread of excitation was perpendicular to the endocardial surface in the central part of the interventricular septum and in the free wall of the left ventricle. In the basal and apical parts of the interventricular septum and in the free wall of the right ventricle excitation spreads in a tangential fashion.

We would like to thank Prof. D. Durrer and Dr. G. Janse from Amsterdam for reviewing the manuscript for the criticism and for their many helpful suggestions.

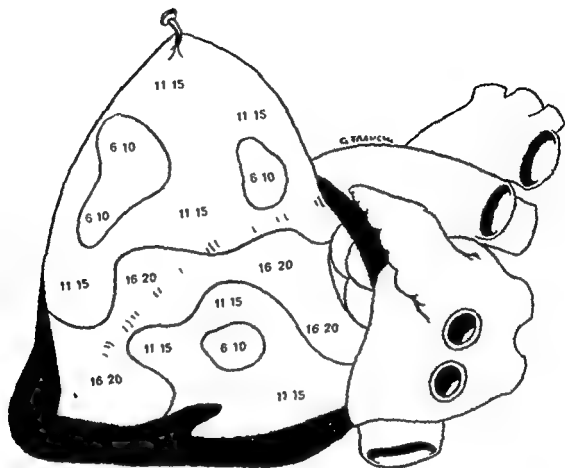


Fig. 41 Endocardial activation of a fetal human heart (subject age 6 $\frac{1}{2}$ months heart weight 5.5 gram). Same heart as in Fig. 34. In this figure an ideal cut has been made along the posterior interventricular septum. Here we see activation of the left endocardium.

concentric isochrones toward the periphery of the septal and parietal endocardium the general direction of activation is from apex toward base and from left to right. The basal endocardium and the septum just below the pulmonic valves are activated last synchronously with the end of QRS complex of the reference lead.

If one compares epicardial and endocardial activation in the same heart it becomes readily evident that endocardial excitation of the left ventricle always precedes excitation of the overlying epicardium by at least 10 msec. On the contrary, in the right ventricle endocardial activity is simultaneous with epicardial activation or may even occur later.

The conclusion follows that in the left ventricular wall the activation front proceeds from endocardium towards epicardium along concentric isochrones while activation of the right ventricular wall must occur mainly in a tangential fashion. A

similar pattern with exclusion of the trabecular region was found in adult hearts with normal right ventricle. Fig. 3 depicts the activation of the interventricular septum. The heart is the same as the one shown in Figs. 41 and 42. In this case seven multipolar needle electrodes had been inserted into the septal mass at different levels. The activation which starts in the left central region proceeds smoothly from the left to right in all subendocardial layers and in most of the lower half of the septum.

Approaching the right endocardium a sudden increase of activation times can be found from point to point indicating slowing of conduction. In most of the upper part of the septum and near the apex on the right side the activation occurs almost simultaneously at different points on the same level. Therefore the pattern followed by the isochrones indicates that the excitation front advances perpendicularly to the septal endocardium in the central portion

- recorded in the course of seven cases of heart surgery *Circulation* 5:48 1952
- 14 Jouve A Corniol J Velasque P Benjamine R and Peytavy R Les dérivationes épiscopales ventriculaires chez l'homme *Acta Cardiol* 11: 247 1958
- 15 Barbato E Pileggi F and Debes A C Study of the sequence of ventricular activation and the QRS complex of the normal human heart using direct epicardial leads *AM HEART J* 5: 867 1958
- 16 Boineau J P Hill J D Spach M S and Moore N The electrocardiogram in right ventricular hypertrophy *AM HEART J* 16: 605 1968
- 17 Brusca A Soleno F and Actis Dato A An electrographic study of right ventricular hypertrophy due to pulmonary stenosis *AM HEART J* 57: 134 1959
- 18 Boineau J P Spach M S and Ayers C R Genesis of the electrocardiogram in atrial septal defect *AM HEART J* 111: 637 1964
- 19 Wallace A G Spach M S Estes M H and Boineau J P Activation of the normal and hypertrophied human right ventricle *AM HEART J* 75: 728 1968
- 20 Roos J P Van Dam R Th and Durrer D Epicardial and intramural excitation of the normal heart in six patients 50 years of age and older *Br Heart J* 30: 630 1968
- 21 Durrer D Buller J Graaff P Lo J G and Meijer F L Epicardial excitation pattern as observed in the isolated revived and perfused human heart *Circ Res* 9: 29 1961
- 22 Durrer D Electrical aspects of human cardiac activity a clinical physiological approach to excitation and stimulation *Cardiovasc Res* 2: 1 1968
- 23 Durrer D Van Dam R Th Freud G E Janse M J Meijer F L and Arzbacher M C Total excitation of the isolated human heart *Circulation* 41: 899 1970
- 24 Alariste V M Cisneros F Diaz de Leon M and Mota A Los indices de Lewis y de Jinich en el recién nacido *Arch Cardiol Mexico* 39: 500 1969
- 25 Moore E N Melbin J Spear J F and Hill J D Sequence of atrial excitation in the dog during antegrade and retrograde activation *J Electrocardiol* 4: 283 1972
- 26 Paes de Carvalho A De Mello W C and Hoffman B F Electrophysiological evidence for specialized fiber types in rabbit atrium *Am J Physiol* 196: 483 1959
- 27 Vaalle M and Hoffman B F Spread of sinus activation during potassium administration *Circ Res* 17: 285 1965
- 28 King T D Barr R C Herman Giddens G S Boaz D M and Spach M S Isopotential body surface maps and their relationship to atrial potentials in the dog *Circ Res* 30: 393 1972
- 29 Amer N S Stuke J H Hoffman B F Cappeletti R R and Domingo R T Activation of the interventricular septal myocardium studied during cardiopulmonary bypass *AM HEART J* 59: 724 1960

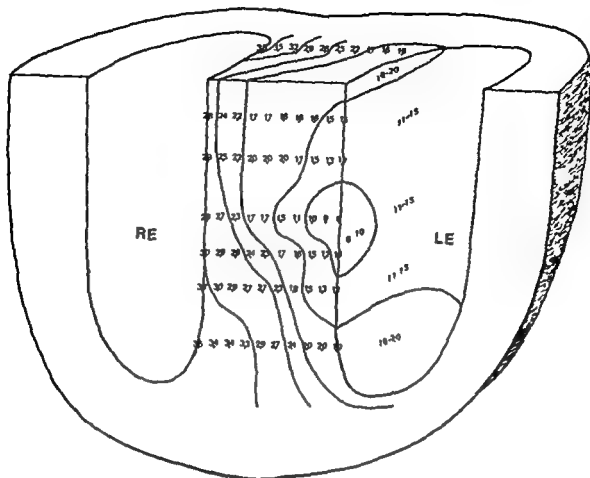


Fig 5 Activation of the interventricular septum LE = left endocardium RE = right endocardium The anterior apical region is not included in the figure

REFERENCES

- 1 Taccardi B Distribution of heart potentials on thoracic surface of normal human subjects *Circ Res* 12:341 1963
- 2 Boineau J and Spach M S The relationship between the electrocardiogram and the electrical activity of the heart *J Electrocardiol* 1:117 1968
- 3 Rosenbloom M R Elizari M V and Lazzari E Los hemibloques Buenos Aires 1967 Ed Paidós
- 4 Lewis T and Rothschild M A The excitatory process in the dog's heart II The ventricles *Phil Trans R Soc (Lond)* 206:181 1915
- 5 Sodi Pallares D New bases of electrocardiography St Louis 1956 The C V Mosby Company
- 6 Scher A M and Young A G Ventricular depolarization and the genesis of QRS *Ann N Y Acad Sci* 65:768 1956 57
- 7 Durrer D Roos J P and Buller J The spread of excitation in canine and human heart in Taccardi B and Marchetti G editors International Symposium on Electrophysiology of the Heart Oxford 1965 Pergamon Press pp 203 214
- 8 Brusca A Bragazzi E Gandolfo S Lavezaro G Politi G Rosettani E and Nosenzo C A tridimensional study of the activation process of the ventricular wall *Acta Cardiol* 21:644 1966
- 9 Hill D J Moore E N and Patterson D F Ventricular epicardial activation studies in experimental and spontaneous right bundle branch block in the dog *Am J Cardiol* 21:737 1968
- 10 Barker P S MacLeod A G and Alexander J The excitatory process observed in the exposed human heart *Am Heart J* 5:720 1930
- 11 Puddu V and Cammirelli C Un cas de ectopie cordis avec une étude électrocardiographique de la progression de l'onde d'excitation *Arch Mal Coeur* 31:861 1938
- 12 Groedel F M and Borchardt P R Direct electrocardiography of the human heart and intrathoracic electrocardiography New York 1948 Brooklyn Medical Press
- 13 Carouso G J Chevalier H A Latscha I and Lenegre J Epicardial electrocardiogram

- recorded in the course of seven cases of heart surgery *Circulation* 5 48 1952
- 14 Jouve A Corriol J Velasque P Benjamine R and Peytavy M Les dérivationes épicardiques ventriculaires chez l'homme *Acta Cardiol* 18 247 1958
- 15 Barbato E Pileggi F and Debes A C Study of the sequence of ventricular activation and the QRS complex of the normal human heart using direct epicardial leads *Am HEART J* 50:857 1958
- 16 Boineau J P Hill J D Spach M S and Moore N The electrocardiogram in right ventricular hypertrophy *Am HEART J* 76 605 1968
- 17 Brusca A Soleno F and Actis Dato A An electrographic study of right ventricular hypertrophy due to pulmonic stenosis *Am HEART J* 57:134 1959
- 18 Boineau J P Spach M S and Ayers C R Genesis of the electrocardiogram in atrial septal defect *Am HEART J* 68 637 1964
- 19 Wallace A G Spach M S Estes E H and Boineau J P Activation of the normal and hypertrophied human right ventricle *Am HEART J* 70:728 1968
- 20 Roos J P Van Dam R Th and Durrer D Epicardial and intramural excitation of the normal heart in six patients 50 years of age and older *Br Heart J* 30 630 1968
- 21 Durrer D Buller J Graaff P Lo I G and Meyler F L Epicardial excitation pattern as observed in the isolated revived and perfused human heart *Circ Res* 9:29 1961
- 22 Durrer D Electrical aspects of human cardiac activity a clinical physiological approach to excitation and stimulation *Cardiovasc Res* 2 1 1968
- 23 Durrer D Van Dam R Th Freud G E Janse W J Meijler F J and Arzbacher H C Total excitation of the isolated human heart *Circulation* 41:890 1970
- 24 Alatriste V M Cisneros F Diaz de Leon M and Mota A Los indices de Lewis y de Junich en el recién nacido *Arch Cardiol Mexico* 39 500 1969
- 25 Moore E N Melbin J Spear J F and Hill J D Sequence of atrial excitation in the dog during antegrade and retrograde activation *J Electrocardiol* 4 283 1972
- 26 Paes de Carvalho A De Mello W C and Hoffman B F Electrophysiological evidence for specialized fiber types in rabbit atrium *Am J Physiol* 196 483 1959
- 27 Vassalle M and Hoffman B F Spread of sinus activation during potassium administration *Circ Res* 17 285 1965
- 28 King T D Barr R C Herman Giddens G S Boaz D E and Spach M S Isopotential body surface maps and their relationship to atrial potentials in the dog *Circ Res* 30 393 1972
- 29 Amer N S Stuke J H Hoffman B F Cappelletti R R and Domingo R T Activation of the interventricular septal myocardium studied during cardiopulmonary bypass *Am HEART J* 59 224 1960

Hemodynamic changes during complete heart block in the unanesthetized monkey

Ralph P Forsyth Ph D
L Henry Edmunds, Jr, M D
David W Amory M D *
Kenneth L Melmon M D
Pale D Thomson, M D
San Francisco Calif

Complete atrioventricular (A V) block produces progressive heart failure and myocardial hypertrophy in dogs and cats^{1,2} and cardiomegaly with or without syncope episodes in man. In man heart failure often accompanies complete heart block, but usually the failure is in part related to additional heart disease. Children with complete heart block generally have higher idioventricular cardiac rates (40 to 80) than do adults (30 to 60). These observations suggest that both age and species affect the circulatory adjustments made after the onset of complete heart block.

In an effort to better understand these mechanisms we studied the changes in regional blood flow that occurred during a six hour period of complete A V block in unanesthetized adult rhesus monkeys with radionuclide labeled microspheres.³ We compared these data with those obtained previously in unanesthetized monkeys during stimulation of the sympathetic nervous system.

Methods

Six male monkeys (*Macaca mulatta*) weighing from 4.0 to 8.0 kilograms were used in this study. During sodium pentobarbital anesthesia (30 mg per kilogram of body weight) polyvinyl catheters were implanted in the abdominal aorta and ventrally in the iliac vessels and in the left ventricle of the heart via the left common carotid artery. The other end of each catheter was brought out through the skin near the umbilicus. After recovery, the monkeys were placed in restraining chairs inside sound protected booths; each catheter was continuously flushed with lightly heparinized (5 units per milliliter) 0.9 percent NaCl at 1 ml per hour.

Three to five days later complete heart block was produced. The animals were anesthetized with sodium pentobarbital and mechanically ventilated with room air. Arterial pressures, heart rate and an electrocardiogram (ECG) were monitored throughout the procedure. After a right

From the Cardiovascular Research Institute, Division of Clinical Pharmacology in Department of Medicine and Department of Surgery, University of California San Francisco, San Francisco, California 94122.

This study was supported in part by NIH Program Project Grants HL 06 85 and GM 16496 and NIH Grants HL 13105 and HL 09964.

Received for publication Aug. 18, 1972.

Reprint requests to: Dr. Ralph P. Forsyth, Cardiovascular Research Institute, University of California San Francisco Medical Center, San Francisco, California 94122.

*Present address: School of Medicine, University of Washington, Seattle, Wash.

anterolateral thoracotomy the pericardium was opened and two insulated pacemaker wires (American Cyanamid Company) were sutured in the body of the right ventricle. While venous inflow was occluded for 15 to 25 minutes a right atriotomy was performed and the bundle of His near the A-V node was destroyed by either ligation or electrocautery. After ECG confirmation of complete A-V dissociation the ventricular pacing electrodes were stimulated at the rate that approximated each animal's unanesthetized base line level. The chest was closed with drainage the pacing wires were passed subcutaneously to the umbilical area and brought out near the catheters. After recovery the monkeys were again placed in a sitting position in their restraining chairs.

The experiments were performed five to eight days following the heart block operation. An hour before base line measurements were made the booth doors were closed to reduce environmental influences the booths were ventilated and lit as during the previous days. Pressure from each catheter was continuously measured with Statham P23 Gb pressure transducers placed at midthoracic levels. Before cardiac output was measured a 2 ml arterial blood sample was taken for measurement of hematocrit, pH, P_{O_2} and P_{CO_2} , the latter measured with Radiometer microelectrodes at 35°C. Cardiac output was measured in duplicate by the indicator dilution method. Indocyanine green was injected into the left ventricle and the concentration of dye in blood withdrawn at a constant rate from the thoric catheter was measured using a Waters X302 densitometer. Measurements were recorded on a type R Beckman recorder.

Directly following the cardiac output measurement distribution of cardiac output was determined with the use of radioactive microspheres. This technique allows up to five independent measurements in each animal.⁴ After base line measurements were obtained the pacemaker was turned off subsequent measurements were made at 15 minutes (second microsphere injection), two hours (third injection) and six hours (fourth injection). Following the fourth injection the pacemaker was restarted (at base line rates) and a fifth set

of measurements was made 30 minutes later.

For each regional blood flow measurement a batch of 50 μ diameter carbonized microspheres labeled with either ^{141}Ce , ^{51}Cr , ^{86}Sr or ^{90}Nb in 10 ml of 0.9 per cent NaCl was injected over a 15 to 20 second period through the left ventricular catheter. Each suspension contained 10,000 to 20,000 microspheres (0.8 to 2 million counts per minute of isotope) which mixed with blood in the left ventricle and was distributed to each organ in direct proportion to organ blood flow. Microspheres are trapped in systemic arterioles. We have estimated that about 0.1 per cent of the monkey's arterioles are occluded by each batch of microspheres.⁴

After the experiment monkeys were killed with an overdose of sodium pentobarbital and all organs and tissues were removed weighed and placed in glass vials. Each vial was counted for four minutes with a Nuclear Chicago scintillation counter (25 to 30 per cent of skeletal muscle limb bone skin chest wall skull and spine were counted all other organs were counted in their entirety). Counts from the various radionuclides were differentiated as described previously.⁴ For each determination the fraction of cardiac output to each organ was calculated from the amount of radioactivity in that organ as compared to the total body count for each isotope. Flow to each organ was determined by multiplying the percentage of cardiac output to that organ by the dye dilution cardiac output. Organ resistance was calculated as the mean driving pressure (mean systemic arterial pressure minus central venous pressure) divided by flow. Details of this procedure and estimates of the reliability of measurements and base line values have been previously reported.^{4,5}

Serial measurements of the same systemic and regional variables and by the same techniques in control monkeys with out heart block remained stable during time periods similar to those of the present study.⁵ The studies with control monkeys also indicated that microsphere infusions do not influence subsequent measurements although total peripheral resistance increases slightly and occasionally transient arrhythmias occur.

Table 1 Hemodynamic measurements before each of the five injections of microspheres for the six experimental monkeys

| Measurement | Microsphere injection | | | | | | | | | |
|--|-----------------------|-----|--------------------|-----|------------------|-----|------------------|-----|------------------------|-----|
| | 1 | | 2 | | 3 | | 4 | | 5 | |
| | Base line | | 15 min after block | | 2 hr after block | | 6 hr after block | | After pacing restarted | |
| | Mean | SD | Mean | SD | Mean | SD | Mean | SD | Mean | SD |
| Systolic arterial pressure (mm Hg) | 129 | 17 | 123 | 19 | 119 | 14 | 124 | 13 | 122 | 3 |
| Diastolic arterial pressure (mm Hg) | 65 | 12 | 46* | 14 | 45* | 11 | 49* | 12 | 71 | 7 |
| Mean arterial pressure (mm Hg) | 95 | 13 | 75* | 17 | 73* | 13 | 80* | 12 | 83 | 14 |
| Cardiac output (ml/min/kg) | 264 | 62 | 188* | 68 | 145* | 86 | 231 | 126 | 304 | 163 |
| Total peripheral resistance (mm Hg/L/min/kg) | 368 | 117 | 424 | 119 | 595* | 209 | 415 | 174 | 351 | 125 |
| Pulse rate (beats/min) | 171 | 13 | 95* | 21 | 88* | 28 | 92* | 33 | 171 | 3 |
| Stroke volume (ml/min/kg) | 1.6 | 0.7 | 2.0* | 0.8 | 1.8 | 0.9 | 2.6* | 2.6 | 1.8 | 1.1 |
| Left ventricular end diastolic pressure (mm Hg)† | 6 | 11 | 12* | 18 | 9 | 12 | 7 | 5 | 5 | 8 |
| Central venous pressure (mm Hg)† | 1 | 8 | 8* | 10 | 4 | 19 | 2 | 5 | 2 | 8 |

*Significantly changed from base line ($P < 0.05$ paired t test)† $n = 4$ measurement values

Results

The overall cardiovascular measurements observed before each of the five microsphere injections are presented in Table I. Changes in the distribution of cardiac output, regional organ blood flows, and resistances are shown in Tables II, III, and IV respectively.

Pulse rate and diastolic and mean systemic arterial pressures fell immediately after stopping the pacemaker. At 15 minutes these measurements and cardiac output were significantly lower than base line. Stroke volume, left ventricular end diastolic, and central venous pressures increased significantly. At this time blood was distributed preferentially toward heart, brain, liver (hepatic artery), skull, spine, and adrenals at the expense of kidneys, skin, skeletal muscle, spleen, and pancreas. Blood vessels in organs that received an increased fraction of cardiac output (except for skull, spine, and adrenals) dilated

vessels to those organs receiving a decreased fraction were constricted.

Two hours after block, systemic arterial pressures and pulse rate were still near those observed at 15 minutes. Cardiac output had fallen further, and total peripheral resistance had increased significantly compared to base line. Left ventricular end diastolic and central venous pressures fell to nearly base line levels and were unchanged at subsequent measurement periods. The pattern of distribution of cardiac output was similar to that found 15 minutes after block, except for the kidneys. Compared to observations at 15 minutes, resistances in each of the measured regional beds showed some increase (except for hepatic artery, pancreas, and adrenals).

Six hours after block, the hemodynamic pattern had changed. Although diastolic arterial pressure and pulse rate remained lower than base line, cardiac output rose due to increased stroke volume and total

1 1 m 11
N mbe 1

Table II Absolute values in the fraction of cardiac output to selected organs in the six experimental monkeys

| | Microsphere injection | | | | | | | | | |
|--------------------------|--|-----|--------------------------|-----|------------------------|-----|------------------------|-----|------------------------|-----|
| | 1 | | 2 | | 3 | | 4 | | 5 | |
| | Absolute fraction of cardiac output at base line | | 15 min after heart block | | 2 hr after heart block | | 6 hr after heart block | | After pacing restarted | |
| | Mean | SD | Mean | SD | Mean | SD | Mean | SD | Mean | SD |
| Heart | 4.2 | 1 | 5.8* | 1 | 5.9 | 1 | 4.3 | 1 | 5.7 | 2 |
| Brain | 6.2 | 2 | 9.4 | 3 | 10.0 | 4 | 9.3* | 3 | 6.6 | 1 |
| Kidney | 17.2 | 2 | 13.9 | 4 | 15.0 | 3 | 14.7 | 2 | 13.6 | 2 |
| Skin | 6.2 | 1 | 3.7 | 1 | 3.9 | 1 | 4.7 | 1 | 4.6 | 2 |
| Skeletal muscle | 23.8 | 4 | 19.0 | 2 | 15.8 | 1 | 15.7 | 3 | 19.4 | 5 |
| Gastrointestinal organs† | 7.0 | 2 | 7.1 | 2 | 7.4 | 2 | 8.5 | 3 | 8.3 | 3 |
| Spleen | 2.6 | 1 | 1.7 | 1 | 1.6 | 1 | 1.6 | 1 | 2.2 | 1 |
| Pancreas | 1.7 | 0.4 | 1.0 | 0.6 | 1.2 | 0.3 | 1.5 | 0.3 | 1.7 | 0.3 |
| Liver (hepatic artery) | 5.9 | 1 | 9.2* | 2 | 12.3 | 3 | 11.5* | 4 | 8.1 | 2 |
| Limb bone | 3.8 | 1 | 4.5 | 2 | 4.9 | 2 | 6.2 | 2 | 5.0 | 2 |
| Skull and spine | 7.9 | 2 | 10.0 | 2 | 9.4 | 2 | 8.9 | 1 | 9.6 | 2 |
| Adrenals | 0.2 | 0.1 | 0.3 | 0.1 | 0.3 | 0.1 | 0.3 | 0.0 | 0.3 | 0.0 |

*Significantly changed from base line ($P < 0.05$ paired t test)
†Cl of stomach, small and large intestine and cecum

peripheral resistance fell to nearly base line levels. Compared to values at two hours all organs except the heart showed some vasodilatation. Blood flow was preferentially shunted to the brain, liver (hepatic artery), limb bone and adrenals at the expense of skin, skeletal muscle and spleen. However, compared to base line measurements, absolute values of blood flow were unchanged except for lowered flow to the kidneys, skeletal muscle, spleen and pancreas.

Thirty minutes after the pacing had resumed, there were no significant differences compared to base line in any of the systemic hemodynamic measurements, regional blood flow or resistances. The fraction of cardiac output at this time was increased to the coronary and hepatic arteries, skull, spine and adrenals, that to the kidney, skin and skeletal muscle remained decreased.

Other organs measured but not included in the tables (eyes, lymph nodes, lung [bronchial artery], chest wall, diaphragm, thyroid and fat) had no significant redistribution of cardiac output; thus their

blood flow and resistance changed in the same proportion as the cardiac output. There were no significant changes in P_{O_2} , P_{CO_2} , pH or hematocrit during the experiment.

The monkeys showed no gross behavioral effects from the experimental procedure. They continued their usual activities and did not struggle or show other signs of discomfort. Gross examination of organs at autopsy revealed no consistent pathological changes, although in two animals portions of subcutaneous tissue along the catheter tracks showed signs of infection. There were no signs of sepsis during life.

Discussion

The base line hemodynamic measurements we obtained prior to turning off the pacemaker were not significantly different from those we have obtained in other monkeys.²⁻⁶ Thus the heart block operation and period of electrical pacing did not alter the measured cardiovascular variables. In comparison to the dog and man, the monkey has a high resting heart rate and cardiac index. However, the percentage

Table III Absolute values for flow (milliliters per 100 Gm) to selected organs in the six experimental monkeys

| | Microsphere injection | | | | | | | | | |
|--------------------------|-----------------------|-----|--------------------------|-----|------------------------|-----|------------------------|-----|------------------------|-----|
| | 1 | | 2 | | 3 | | 4 | | 5 | |
| | Base line | | 15 min after heart block | | 2 hr after heart block | | 6 hr after heart block | | After pacing restarted | |
| | Mean | SD | Mean | SD | Mean | SD | Mean | SD | Mean | SD |
| Heart | 317 | 75 | 270 | 82 | 223* | 98 | 278 | 77 | 432 | 186 |
| Brain | 98 | 37 | 95 | 30 | 77 | 27 | 124 | 57 | 117 | 67 |
| Kidney | 1138 | 286 | 634* | 247 | 614* | 211 | 910* | 293 | 1044 | 283 |
| Skin | 24 | 8 | 9* | 5 | 9* | 6 | 18 | 12 | 20 | 17 |
| Skeletal muscle | 21 | 7 | 12* | 6 | 8* | 5 | 12* | 7 | 20 | 13 |
| Gastrointestinal organs† | 72 | 24 | 47* | 17 | 42* | 19 | 72 | 34 | 96 | 52 |
| Spleen | 425 | 49 | 210* | 51 | 168* | 74 | 268* | 57 | 389 | 73 |
| Pancreas | 320 | 126 | 131* | 77 | 137* | 81 | 224* | 110 | 348 | 103 |
| Liver (hepatic artery) | 54 | 14 | 55 | 18 | 65 | 40 | 90 | 54 | 84 | 54 |
| Lungs (bronchial artery) | 62 | 56 | 45 | 38 | 48 | 27 | 64 | 50 | 79 | 55 |
| Limb bone | 9 | 3 | 8 | 5 | 8 | 9 | 14 | 12 | 16 | 15 |
| Skull and spine | 22 | 9 | 19 | 6 | 15 | 9 | 22 | 12 | 30 | 18 |
| Adrenals | 227 | 71 | 211 | 132 | 201 | 115 | 276 | 125 | 342 | 184 |

*Significantly changed from base line ($P < 0.05$ paired t test)

†Includes stomach, small and large intestine, and cecum

decreases in heart rate and cardiac output that we found after electrical pacing was stopped were similar to those in dogs.^{2,3,7,9} Our data also show that the monkey like the dog, has an increased pulse pressure, total peripheral resistance, stroke volume, left ventricular end diastolic and central venous pressures during complete heart block. We did not observe the six to twelve seconds of asystole that White and associates⁹ reported in dogs immediately after cessation of pacing.

The increase of cardiac output and stroke volume six hours after the onset of heart block suggests that compensatory influences (such as sodium retention and an increased blood volume), may have begun. Compensatory adjustments have been reported much later after A-V block.^{1,2} Central venous and left ventricular end diastolic pressures were near base line values six hours after block, although diastolic and mean systemic arterial pressures were still significantly lower.

Except for coronary flow measurements there are few data about the redistribution of cardiac output after onset of complete

heart block. In unanesthetized dogs⁸ the onset of heart block decreased iliac arterial blood flow more than circumflex coronary and mesenteric blood flows and did not change renal flow. In unanesthetized monkeys blood flow to skeletal muscle and gastrointestinal organs decreased during heart block but renal blood flow also decreased and did not begin to increase until six hours after heart block.

Several reports have noted significant decreases in coronary blood flow in dogs after experimental heart block. In anesthetized dogs coronary blood flow was found to be reduced to 50 per cent of preblock values 60 to 90 minutes after section of the bundle of His.⁷ In the unanesthetized dog White and associates⁹ reported that circumflex coronary blood flow decreased 37 per cent a few minutes after cessation of pacing. Pitt and Gregg¹⁰ found similar reductions in coronary flow when their dogs were paced at a mean rate of 63 times per minute compared to a mean rate of 152 times per minute. In our monkeys coronary blood flow decreased less, falling an average of 15 per cent at

Table IV Changes in regional resistances expressed as a per cent of base line values in the six experimental monkeys

| | Microsphere injection | | | | | | | |
|--------------------------|--------------------------|------|------------------------|------|------------------------|------|------------------------|------|
| | 2 | | 3 | | 4 | | 5 | |
| | 15 min after heart block | | 2 hr after heart block | | 6 hr after heart block | | After pacing restarted | |
| | Mean | S.D. | Mean | S.D. | Mean | S.D. | Mean | S.D. |
| Heart | 87 | 34 | 112 | 39 | 109 | 54 | 80 | 30 |
| Brain | 80 | 22 | 101 | 31 | 57 | 19 | 107 | 86 |
| Kidney | 150 | 24 | 192 | 58 | 114 | 36 | 130 | 77 |
| Spleen | 204 | 40 | 260 | 48 | 126 | 41 | 147 | 70 |
| Skin | 145 | 24 | 234* | 65 | 158 | 45 | 134 | 84 |
| Skeletal muscle | 121 | 25 | 150 | 63 | 95 | 24 | 88 | 47 |
| Gastrointestinal organs† | 186 | 83 | 363 | 133 | 127 | 47 | 127 | 51 |
| Liver | 246 | 59 | 212 | 87 | 115 | 49 | 110 | 68 |
| Pancreas | 81 | 21 | 81 | 37 | 60 | 37 | 82 | 50 |
| Lung (hepatic artery) | 187 | 85 | 126 | 109 | 131 | 76 | 117 | 93 |
| Lung (bronchial artery) | 112 | 50 | 153 | 66 | 66 | 40 | 110 | 107 |
| Limb bone | 95 | 29 | 134 | 57 | 72 | 25 | 90 | 61 |
| Skull and spine | 103 | 53 | 93 | 47 | 64 | 22 | 74 | 33 |
| Adrenals | | | | | | | | |

*Significantly changed from base line ($P < 0.05$ paired t test)
 †Included stomach, small and large intestine and cecum

15 minutes after block and 30 per cent two hours after block. At both these times the coronary vascular bed received an increased fraction of cardiac output, coronary flow decreasing less than the cardiac output.

Aside from the species difference and the method of creating heart block, measurement technique is the most important factor in the interpretation of these discrepant findings. Radioactive microspheres do not measure flow continuously, but do estimate blood flow without interfering with vessels to major organs. Previous studies using unanesthetized dogs^{8,10} estimated total coronary flow with previously placed flow meters on the circumflex branch of the left coronary artery. This technique requires extensive surgical manipulation, may cause partial occlusion of the artery, and assumes that circumflex blood flow represents total coronary flow.

Patients with chronic complete heart block and associated myocardial disease have a reduced blood flow to the heart¹¹ and brain.¹² In these studies restoration of normal heart rate by pacing increased

blood flow to both brain and heart. Our data in monkeys indicate that neither the heart (except at the second hour observation) nor the brain was underperfused during six hours of idioventricular rhythm and that subsequent pacing increased blood flow to the heart but not to the brain. These human studies cannot be directly compared to our primate results since older patients with complete heart block generally have extensive coronary atherosclerosis. Our observations in adult monkeys may be more comparable to acute heart block in children. These patients usually have higher idioventricular rates, are often asymptomatic, and do not have advanced atherosclerosis.

The unanesthetized primate preserves blood flow to the heart, brain, liver (hepatic artery), bone, and adrenals after onset of complete heart block by decreasing blood flow to skeletal muscle, gastrointestinal organs, and kidneys. Six hours after onset of block, cardiac output increased toward base line, but blood flow to skeletal muscle, kidneys, spleen, and pancreas was still decreased. These data suggest that

Table III Absolute values for flow (milliliters per 100 Gm) to selected organs in the six experimental monkeys

| | Microsphere injection | | | | | | | | | |
|--------------------------|-----------------------|-----|--------------------------|-----|------------------------|-----|------------------------|-----|------------------------|-----|
| | 1 | | 2 | | 3 | | 4 | | 5 | |
| | Base line | | 15 min after heart block | | 2 hr after heart block | | 6 hr after heart block | | After pacing restarted | |
| | Mean | S D | Mean | S D | Mean | S D | Mean | S D | Mean | S D |
| Heart | 317 | 75 | 270 | 82 | 223* | 98 | 278 | 77 | 432 | 186 |
| Brain | 98 | 37 | 95 | 30 | 77 | 27 | 124 | 57 | 117 | 67 |
| Kidney | 1138 | 286 | 634* | 247 | 614* | 211 | 910* | 293 | 1044 | 288 |
| Skin | 24 | 8 | 9* | 5 | 9* | 8 | 18 | 12 | 20 | 17 |
| Skeletal muscle | 21 | 7 | 12* | 6 | 8* | 5 | 12* | 7 | 20 | 13 |
| Gastrointestinal organs† | 72 | 24 | 47* | 17 | 42* | 19 | 72 | 34 | 96 | 57 |
| Spleen | 425 | 49 | 210* | 51 | 168* | 74 | 268* | 57 | 389 | 73 |
| Pancreas | 320 | 126 | 131* | 77 | 137* | 81 | 224* | 110 | 348 | 103 |
| Liver (hepatic artery) | 54 | 14 | 55 | 18 | 65 | 40 | 90 | 54 | 84 | 54 |
| Lungs (bronchial artery) | 62 | 56 | 45 | 38 | 48 | 27 | 64 | 50 | 79 | 55 |
| Limb bone | 9 | 3 | 8 | 5 | 8 | 9 | 14 | 12 | 16 | 15 |
| Skull and spine | 22 | 9 | 19 | 6 | 15 | 9 | 22 | 12 | 30 | 18 |
| Adrenals | 227 | 71 | 211 | 132 | 201 | 115 | 276 | 125 | 342 | 184 |

*Significantly changed from base line ($P < 0.05$ paired t test)

†Includes stomach, small and large intestine, and cecum

decreases in heart rate and cardiac output that we found after electrical pacing was stopped were similar to those in dogs.^{2,7,8} Our data also show that the monkey like the dog has an increased pulse pressure, total peripheral resistance, stroke volume, left ventricular end diastolic and central venous pressures during complete heart block. We did not observe the six to twelve seconds of asystole that White and associates⁸ reported in dogs immediately after cessation of pacing.

The increase of cardiac output and stroke volume six hours after the onset of heart block suggests that compensatory influences (such as sodium retention and an increased blood volume), may have begun. Compensatory adjustments have been reported much later after A-V block.^{1,2} Central venous and left ventricular end diastolic pressures were near base line values six hours after block, although diastolic and mean systemic arterial pressures were still significantly lower.

Except for coronary flow measurements, there are few data about the redistribution of cardiac output after onset of complete

heart block. In unanesthetized dogs⁸ the onset of heart block decreased iliac arterial blood flow more than circumflex coronary and mesenteric blood flows and did not change renal flow. In unanesthetized monkeys blood flow to skeletal muscle and gastrointestinal organs decreased during heart block but renal blood flow also decreased and did not begin to increase until six hours after heart block.

Several reports have noted significant decreases in coronary blood flow in dogs after experimental heart block. In anesthetized dogs, coronary blood flow was found to be reduced to 50 per cent of preblock values 60 to 90 minutes after section of the bundle of His.⁷ In the unanesthetized dog White and associates⁸ reported that circumflex coronary blood flow decreased 37 per cent a few minutes after cessation of pacing. Pitt and Gregg¹⁰ found similar reductions in coronary flow when their dogs were paced at a mean rate of 63 times per minute compared to a mean rate of 152 times per minute. In our monkeys coronary blood flow decreased less, falling an average of 15 per cent at

17 mcs 86
N mbr 3

Table IV Changes in regional resistances expressed as a per cent of base line value in the six experimental monkeys

| | Microsphere injection | | | | | | | |
|--------------------------|----------------------------------|------|--------------------------------|------|--------------------------------|------|--------------------------------|------|
| | 2 15 min after heart block | | 3 2 hr after heart block | | 4 6 hr after heart block | | 5 After pacing restarted | |
| | Mean | S.D. | Mean | S.D. | Mean | S.D. | Mean | S.D. |
| Heart | 87 | 34 | 112 | 39 | 109 | 54 | 80 | 30 |
| Brain | 80 | 22 | 101 | 31 | 57 | 19 | 107 | 26 |
| Kidney | 150 | 24 | 192 | 38 | 114 | 36 | 130 | 77 |
| Skin | 204 | 40 | 260 | 48 | 126 | 41 | 147 | 70 |
| Skeletal muscle | 145* | 34 | 234 | 65 | 158 | 45 | 134 | 84 |
| Gastrointestinal organs† | 171 | 25 | 150 | 63 | 95 | 24 | 88 | 47 |
| Spleen | 186 | 83 | 365 | 133 | 127 | 47 | 122 | 51 |
| Pancreas | 246 | 59 | 212 | 87 | 115 | 40 | 110 | 68 |
| Liver (hepatic artery) | 81 | 77 | 81 | 37 | 60 | 37 | 87 | 50 |
| Lung (bronchial artery) | 187 | 83 | 126 | 109 | 131 | 76 | 117 | 91 |
| Limb bone | 112 | 50 | 153 | 66 | 66 | 40 | 110 | 107 |
| Skull and spine | 95 | 29 | 151 | 37 | 72 | 23 | 90 | 61 |
| Adrenals | 103 | 33 | 43 | 47 | 64 | 22 | 74 | 35 |

S.D. = standard deviation from base line ($P < 0.05$ paired t test)
† Included stomach, small and large intestine and rectum

15 minutes after block and 30 per cent two hours after block. At both these times the coronary vascular bed received an increased fraction of cardiac output, coronary flow decreasing less than the cardiac output.

Aside from the species difference and the method of creating heart block, measurement technique is the most important factor in the interpretation of these discrepant findings. Radioactive microspheres do not measure flow continuously but do estimate blood flow without interfering with vessels to major organs. Previous studies using unanesthetized dogs¹⁴ estimated total coronary flow with previously placed flow meters on the circumflex branch of the left coronary artery. This technique requires extensive surgical manipulation, may cause partial occlusion of the artery, and assumes that circumflex blood flow represents total coronary flow.

Patients with chronic complete heart block and associated myocardial disease have a reduced blood flow to the heart¹¹ and brain.¹² In these studies restoration of normal heart rate by pacing increased

blood flow to both brain and heart. Our data in monkeys indicate that neither the heart (except at the second hour observation) nor the brain was underperfused during six hours of idioventricular rhythm and that subsequent pacing increased blood flow to the heart but not to the brain. These human studies cannot be directly compared to our primate results since older patients with complete heart block generally have extensive coronary atherosclerosis. Our observations in adult monkeys may be more comparable to acute heart block in children. These patients usually have higher idioventricular rates, are often asymptomatic, and do not have advanced atherosclerosis.

The unanesthetized primate preserves blood flow to the heart, brain, liver (hepatic artery), bone, and adrenals after onset of complete heart block by decreasing blood flow to skeletal muscle, gastrointestinal organs, and kidneys. Six hours after onset of block, cardiac output increased toward base line, but blood flow to skeletal muscle, kidneys, spleen, and pancreas was still decreased. These data suggest that

blood is redistributed after onset of acute heart block until compensatory changes in blood volume are completed and until cardiac output increases toward normal. Although we did not measure blood volume or urinary sodium levels, our data indicate that compensatory mechanisms increase cardiac output in monkeys within a few hours after the onset of heart block.

The changes in the distribution of cardiac output during acute heart block were similar to those that we have observed in unanesthetized monkeys during hemorrhage.¹³ In both these experimental conditions the fraction of cardiac output to heart, brain, liver (hepatic artery), and adrenals increased at the expense of spleen, pancreas, kidney, skin and muscle. With the same preparation we have found a similar pattern of redistribution of blood flow during acute psychological stress¹⁴ and during electrical stimulation of pressor areas of the hypothalamus.¹⁵ All these data suggest that at least some of the compensatory adjustments to the bradycardia and hypotension caused by A-V block are mediated by an increased discharge from the sympathetic nervous system. Gaffney and Braunwald¹⁶ have noted that the adrenergic nervous system plays an important role in the circulatory adjustment of patients with congestive heart failure and that anesthesia may dangerously decrease the sympathetic response that preserves blood flow to vital organs.

These experiments indicate that acute heart block in primates without coronary artery disease decreases systemic blood pressure and cardiac output. At the same time blood is redistributed to vital organs (heart and brain) at the expense of less important tissues until compensatory mechanisms restore cardiac output toward normal.

Summary

After surgically produced complete heart block, the hearts of six monkeys were paced electrically for five to eight days at their preoperative heart rates. Simultaneous measurements of regional blood flow and other hemodynamic variables were then obtained before and 15 minutes, two hours, and six hours after stopping electrical pacing and 30 minutes after restarting

pacing. Fifteen minutes after stopping pacing heart rate, cardiac output, and systemic mean and diastolic pressures decreased and stroke volume increased. This pattern persisted throughout the six-hour period of idioventricular rhythm except that left ventricular end diastolic and central venous pressures were increased only at 15 minutes and cardiac output was not significantly decreased at the six-hour period. During heart block, the fraction of cardiac output received by the heart (except at six hours), brain, liver (hepatic artery), skull spine, and adrenals was increased at the expense of skin, kidneys (initially), skeletal muscle, spleen, and pancreas. This pattern of regional redistribution of blood flow is similar to that found during hemorrhage and other conditions in which the sympathetic nervous system is stimulated. After pacing was restarted, all the systemic variables measured returned to base line levels, and the regional blood flows measured did not differ from those observed before the heart block.

We thank Mr. David Tyree and Mrs. Diane Barmby for their assistance in this work.

REFERENCES

- Spann J F Jr, Buccino R A, Sonnenblick E H and Braunwald E. Contractile state of cardiac muscle obtained from cats with experimentally produced ventricular hypertrophy and heart failure. *Circ Res* 21:341 1967.
- Starzl T E and Gaertner R A. Chronic heart block in dogs: A method for producing experimental heart failure. *Circulation* 12:159 1955.
- Turna M, Babotai I, Bussmann W H and Krayenbuhl H P. Haemodynamics of acute and chronic atrioventricular block in dogs. *Cardiovasc Res* 3:209 1969.
- Rudolph A M and Heymann M A. The circulation of the fetus in utero: Methods for studying distribution of blood flow, cardiac output and organ blood flow. *Circ Res* 21:163 1967.
- Hoffbrand V I and Forsyth R P. Validity studies of the radioactive microsphere method for the study of the distribution of cardiac output, organ blood flow and resistance in the conscious rhesus monkey. *Cardiovasc Res* 3:426 1969.
- Forsyth R P, Nies A S, Wyler F, Neutze J M and Melmon K I. The normal distribution of cardiac output in the unanesthetized restrained rhesus monkey. *J Appl Physiol* 25:736 1968.
- Starzl T E, Gaertner R A and Baker R R.

- Acute complete heart block in dogs *Circulation* 12:111 1955
- 8 White S Patrick T Higgins C B Vatner S F Franklin D and Braunwald E Effects of altering ventricular rate on blood flow distribution in conscious dogs *Am J Physiol* 221:1407 1971
- 9 Miller D E Gleason W L Whalen R F Morris J J and McIntosh H D Effect of ventricular rate on the cardiac output in the dog with chronic heart block *Circ Res* 10:658 1962
- 10 Pitt B and Gregg D E Coronary hemodynamic effects of increasing ventricular rate in the unanesthetized dog *Circ Res* 22:753 1968
- 11 Rowe G G Stenlund R R Thomsen J H Terry W and Quermit A S Coronary and systemic hemodynamic effects of cardiac pacing in man with complete heart block *Circulation* 40:839 1969
- 12 Shapiro W and Chawla N P S Observations on the regulation of cerebral blood flow in complete heart block *Circulation* 40:863 1969
- 13 Forsyth H P Hoffbrand B J and Melmon H I Redistribution of cardiac output during hemorrhage in the unanesthetized monkey *Circ Res* 27:111 1970
- 14 Forsyth R P Regional blood flow changes during 72 hour avoidance schedules in the monkey *Science* 173:546 1971
- 15 Forsyth R P Hypothalamic control of the distribution of cardiac output in the unanesthetized rhesus monkey *Circ Res* 26:783 1970
- 16 Gaffney T E and Braunwald E Importance of the adrenergic nervous system in the support of circulatory function in patients with congestive heart failure *Am J Med* 31:320 1963

A theoretical approach to the volume pulse wave

Ryu Nakayama MD*

Toshiyuki Kobayashi, MD*

Kiyoyuki Kimura, MD*

Takehiko Azuma MD**

Tokyo, Japan

A number of works concerning plethysmography have been done since the early description of the plethysmograph in 1732.¹ The method mainly used has been to obtain information about blood flow through a segment.² The segmental blood flow has been evaluated by venous occlusion plethysmography,^{3,4,6,7,9,11,12} which was originally described by Brodie and Russell¹² in 1905. Until recently the value of the venous occlusion plethysmography has been retained as the most reliable method for measuring the rate of total arterial blood flow to the extremity or to a segment of a limb.¹³ The limitations of the method however are considerable.¹⁴ The greatest theoretical disadvantage of venous occlusion is probably that it may be productive of unaccountable physiological changes in venous, arterial and lymph flow dynamics.¹⁵ Therefore the plethysmographer has attempted to analyze a single segmental pulse curve for unrecorded inflow and outflow during a pulse cycle without venous or arterial occlusion. No method up to now has satisfied the experts, due to a lack of precise quantitative information concerning the volume pulse wave.

It is believed that the volume pulse wave is the time course of the difference (D) between volume of inflow (I) and volume of outflow (O).¹⁶ This theory that may be called volume difference theory has been strongly supported. It may be represented by the following equation

$$D = I - O \quad (1)$$

Equation (1) means that the amplitude of the volume pulse wave recorded by the plethysmograph for a given moment equals the subtracted value of the volume of outflow from the volume of inflow at the same moment. This is most fundamental to the utilization of the plethysmograph as a tool in clinical medicine. This is however too simple for explaining everything about the volume pulse wave. The weakest point of the expression is that one may consider that the volume pulse wave indicates flow volume itself though it is expressed as the difference between one flow volume and another.

Method

The digital volume pulse was recorded by an Elema Schönder EMT 510 pneumo-

From the Department of Internal Medicine, National Cancer Center Hospital, Tokyo, Japan.
Received for publication on Aug. 22, 1972.

Reprint requests to: Ryu Nakayama, MD, National Cancer Center Hospital, Department of Internal Medicine, 5-1-1 Tsukiji, Chuo-ku, Tokyo, Japan.

*Department of Internal Medicine, National Cancer Center Hospital, Tokyo, Japan.

**Department of Physiology, Shinshu University Medical School, Matsumoto, Japan.

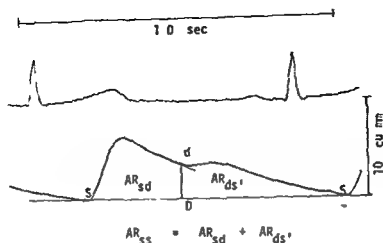


Fig 1 The pattern of volume pulse and the abbreviation of the glossaries S = the starting point of the pulse wave under observation S' = the end of the pulse wave under observation and also the starting point of the successive pulse wave d = the dicrotic notch SD = the time interval from S to d AR_{sd} = the area under the pulse curve over SD interval $AR_{ds'}$ = the area under the pulse curve over DS' interval

plethysmograph connected to a Mingograf Cardirex 6T. For sensitivity standardization a volume calibration was carried out by pushing the calibration button provided in the EMT 510 to a volume that was adjusted to 10 mm³. The subjects involved in this study were placed in two groups. Group A was composed of 40 healthy adults whose ages ranged from 34 to 81 years old and Group B was composed of 24 adults with cardiac disease whose ages ranged from 18 to 80 years old. Group B included 11 cases with cardiac failure. The subjects rested supine in bed in a comfortable environment. The digits studied were usually placed at heart level and sometimes above or below heart level for special observations. In order to observe the effect of venous occlusion on the pulse wave form the brachial vein was occluded by inflation of the cuff wrapped about the arm for several seconds. The plethysmographic cup was placed on the tip of the finger defined as that portion of the finger distal near the terminal interphalangeal articulation. The pattern of volume pulse and abbreviation of glossaries are shown in Figs 1 and 2. Areas under the pulse curve over SD , SS and DS' are named AR_{sd} , AR_{ss} and $AR_{ds'}$ respectively (Fig 1). The areas were measured graphically. A tangent line was drawn along the steepest part of the ascending limb of the pulse curve (Fig 2). The intersections of this line

with the base line and the apex line are named a and b . The time interval between a and b is expressed by Δt which is the so-called inclination time. The height of b point from the base line is expressed by Δt which is equal to the maximum amplitude of the pulse wave. The area under the pulse curve over SB is shown by AR_s . One may draw a line which is parallel to the base line and

$$\frac{AR_s}{SB}$$

apart from the base line. The intersection of this line with the steepest part of the pulse curve is denoted as p . The amplitude of the pulse wave at p is expressed by Δt_p .

Results

As shown in Fig 1 the whole area of the pulse wave (AR) can be divided into two parts - i.e. AR_{sd} and $AR_{ds'}$. When one looks at these areas one may notice that the size of $AR_{ds'}$ correlates to that of AR_{sd} regardless of whether the size of the pulse wave is great or small. Therefore a normal ratio of $AR_{ds'}$ to AR_{sd} could be obtained from Group A and an abnormal ratio if present from Group B. It should be noticed however that the ratio

$$\frac{AR_{ds'}}{AR_{sd}}$$

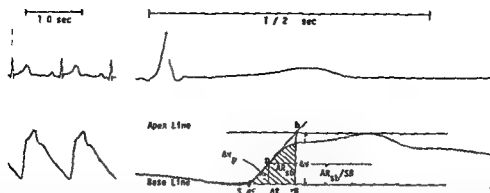


Fig 2 The abbreviation of glossaries $dv/dt = A$ tangent is drawn to the steepest part of the ascending limb of the pulse wave. The time interval between the intersection of this line with the base line (a) and the apex line (b) makes up the inclination time (Δt).²³ The maximum amplitude of the volume pulse is equal to ΔAR_{sb} is the area under the pulse curve over the SB interval. A parallel line to the base line which is AR_{sb}/SB apart from the base line can be drawn. The intersection of this line with the steepest part of the pulse curve is denoted as p. The rate of rise of the pulse wave at the point p is given by

$$\lim_{\Delta t \rightarrow 0} \Delta v / \Delta t = dv/dt$$

SB = the time interval from S to b AR_{sb} = the area under the pulse curve over the time interval between S and b

being within normal limits might be based on the normal ratio

$$\frac{SD}{DS}$$

and that the abnormal ratio

$$\frac{AR_M}{AR_d}$$

might be based on the abnormal ratio

$$\frac{SD}{DS'}$$

though the ratio

$$\frac{AR_d}{AR_d}$$

may have more information than that of the ratio

$$\frac{SD}{DS}$$

In order to find the clinical meaning which the ratios

$$\frac{AR_d}{AR_d}$$

and

$$\frac{AR_d}{AR}$$

might have the values of

$$\frac{AR_M}{SD}, \frac{AR_d}{DS}$$

and

$$\frac{AR}{SS}$$

were calculated. They also closely correlate with each other as shown in Figs 3 and 4. The mean values of the ratio

$$\frac{AR_M}{SD} \cdot \frac{AR_d}{DS}$$

instead of

$$\frac{AR_M}{AP_d}$$

and of the ratio

$$\frac{AR_d}{SD} \cdot \frac{AR}{SS}$$

instead of

$$\frac{AR_d}{AR}$$

were obtained from Group A and B (Table I). The differences between the mean values obtained from Group A and that from Group B were statistically significant.

The correlation between

$$dv/dt$$

and

$$\frac{AR_{sb}}{SB}$$

defined in Fig 2 can be seen in Fig 5

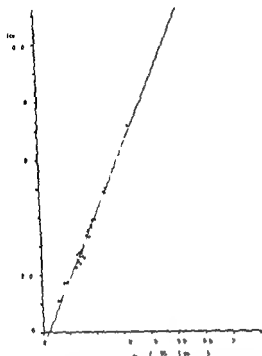


Fig 3 Correlation between AR_m/SD (Y axis) and AR_m/DS' (X axis) in 93 pulse waves obtained from normal subjects. $r = 0.94$ $P < 0.00005$ regression line $y = 2.2x - 0.53$

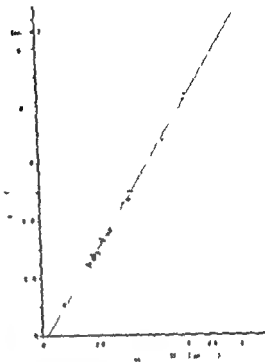


Fig 4 Correlation between AR_m/SD (Y axis) and AR/SS (X axis) in 93 pulse waves obtained from normal subjects. $r = 0.93$ $P < 0.00005$ regression line $y = 1.6x - 0.58$

Changes of the amplitude of the volume pulse wave and the ratio

$$\frac{d/D}{AR/SS}$$

by venous occlusion are shown in Figs 6 and 7 and their mean values are listed in Table II. The difference between the mean value of control and that of venous occlusion was statistically significant in Group A ($P < 0.00005$) but it was not in Group B ($P < 0.1$).

Discussion

At the beginning of the discussion let us try to interpret the actual changes of the pulse wave due to venous occlusion and postural alteration of the limb according to the volume difference theory.

The venous occlusion plethysmographic method of measuring the segmental blood flow is based on the assumption that the initial rate of swelling of the part which indicates the rate of arterial inflow during

venous occlusion is the same as the actual undisturbed rate of arterial inflow immediately before the venous occlusion. If the assumption is accepted it can be thought that the arterial inflow to the digit (I) must be kept unchanged immediately after venous occlusion. Venous drainage (O) is obviously decreased by the occlusion. Therefore volume difference (D) should be increased by the occlusion while the amplitude of the pulse wave actually becomes decreased (Fig 6).

When the digit studied is raised above the heart the amplitude of the volume pulse is actually increased.^{11,13} According to the volume difference theory the size of volume pulsation depends directly on the quantity of the volume difference (D). If the quantity of the volume difference is large enough in comparison with the volume of blood flow, the blood must be progressively accumulating in some part of the vessels. If the quantity of the volume difference is very small in comparison with the volume of blood flow, the volume difference

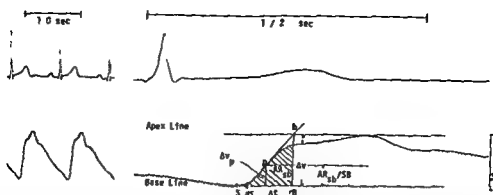


Fig 2 The abbreviation of glossaries dv/dt = A tangent is drawn to the steepest part of the ascending limb of the pulse wave. The time interval between the intersection of this line with the base line (a) and the apex line (b) makes up the inclination time (Δt).¹³ The maximum amplitude of the volume pulse = equal to Δv . AR_{sb} = the area under the pulse curve over the SB interval. A parallel line to the base line which is AR_{sb}/SB apart from the base line can be drawn. The intersection of this line with the steepest part of the pulse curve is denoted as p . The rate of rise of the pulse wave at the point p = given by

$$\lim_{\Delta t \rightarrow 0} \Delta v / \Delta t = dv/dt$$

SB = the time interval from S to B . AR_{sb} = the area under the pulse curve over the time interval between S and B .

being within normal limits might be based on the normal ratio

$$\frac{SD}{DS}$$

and that the abnormal ratio

$$\frac{AR_{sd}}{AR_d}$$

might be based on the abnormal ratio

$$\frac{SD}{DS}$$

though the ratio

$$\frac{AR_d}{AR_d}$$

may have more information than that of the ratio

$$\frac{SD}{DS}$$

In order to find the clinical meaning which the ratios

$$\frac{AR_{sd}}{AR_d}$$

and

$$\frac{AR_{sd}}{AR}$$

might have the values of

$$\frac{AR_d}{SD}, \frac{AR_{sd}}{DS}$$

and

$$\frac{AR}{SS}$$

were calculated. They also closely correlate with each other as shown in Figs 3 and 4. The mean values of the ratio

$$\frac{AR_{sd}}{SD} \cdot \frac{AR_d}{DS}$$

instead of

$$\frac{AR_{sd}}{AR_d}$$

and of the ratio

$$\frac{AR_d}{SD} \cdot \frac{AR}{SS}$$

instead of

$$\frac{AR_d}{AR}$$

were obtained from Group A and B (Table 1). The differences between the mean values obtained from Group A and that from Group B were statistically significant.

The correlation between

$$dv/dt$$

and

$$\frac{AR_s}{SB}$$

defined in Fig 2 can be seen in Fig 5.

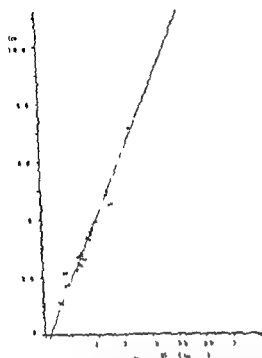
1 me 86
m r 2

Fig 3 Correlation between AR_u/SD (Y axis) and AR_u/DS (X axis) in 93 pulse waves obtained from normal subjects $r = 0.94$ $P < 0.00005$ regression line $Y = 2.2 X - 0.53$

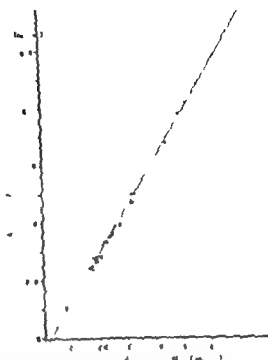


Fig 4 Correlation between AR_u/SD (Y axis) and AR_u/SS (X axis) in 93 pulse waves obtained from normal subjects $r = 0.98$ $P < 0.0000$ regression line $Y = 1.6 X - 0.58$

Changes of the amplitude of the volume pulse wave and the ratio

$$\frac{dV/dt}{\frac{AR_u}{SB}}$$

by venous occlusion are shown in Figs 6 and 7 and their mean values are listed in Table II. The difference between the mean value of control and that of venous occlusion was statistically significant in Group A ($P < 0.00005$) but it was not in Group B ($P < 0.1$).

Discussion

At the beginning of the discussion let us try to interpret the actual changes of the pulse wave due to venous occlusion and point out an alteration of the limb according to the volume difference theory.

The venous occlusion plethysmographic method of measuring the segmental blood flow is based on the assumption that the initial rate of swelling of the part which indicates the rate of arterial inflow during

venous occlusion is the same as the actual undisturbed rate of arterial inflow immediately before the venous occlusion. If the assumption is accepted it can be thought that the arterial inflow to the digit (I) must be kept unchanged immediately after venous occlusion. Venous drainage (O) is obviously decreased by the occlusion. Therefore volume difference (D) should be increased by the occlusion while the amplitude of the pulse wave actually becomes decreased (Fig 6).

When the digit studied is raised above the heart the amplitude of the volume pulse is actually increased.^{11,12} According to the volume difference theory, the size of volume pulsation depends directly on the quantity of the volume difference (D_v). If the quantity of the volume difference is large enough in comparison with the volume of blood flow, the blood must be progressively accumulating in some part of the vessels. If the quantity of the volume difference is very small in comparison with the volume of blood flow, the volume difference

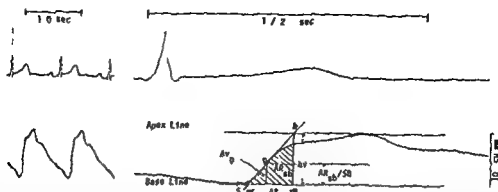


Fig 2 The abbreviation of glossaries ds/dt = A tangent is drawn to the steepest part of the ascending limb of the pulse wave. The time interval between the intersection of this line with the base line (a) and the apex line (b) makes up the inclination time (Δt).¹⁰ The maximum amplitude of the volume pulse is equal to Δs . AR_s = the area under the pulse curve over the SB interval. A parallel line to the base line which is AR_s/SB apart from the base line can be drawn. The intersection of this line with the steepest part of the pulse curve is denoted as p. The rate of rise of the pulse wave at the point p is given by

$$\lim_{\Delta t \rightarrow 0} \frac{\Delta s}{\Delta t} = ds/dt$$

SB = the time interval from S to b. AR_s = the area under the pulse curve over the time interval between S and b.

being within normal limits might be based on the normal ratio

$$\frac{SD}{DS}$$

and that the abnormal ratio

$$\frac{AR_{sd}}{AR_{ds}}$$

might be based on the abnormal ratio

$$\frac{SD}{DS'}$$

though the ratio

$$\frac{AR_d}{AR_s}$$

may have more information than that of the ratio

$$\frac{SD}{DS'}$$

In order to find the clinical meaning which the ratios

$$\frac{AR_d}{AR_s}$$

and

$$\frac{AR_d}{AR}$$

might have the values of

$$\frac{AR_{sd}}{SD} \frac{AR_d}{DS}$$

and

$$\frac{AR}{SS}$$

were calculated. They also closely correlate with each other as shown in Figs 3 and 4. The mean values of the ratio

$$\frac{AR_{sd}}{SD} \frac{AR_d}{DS'}$$

instead of

$$\frac{AR_d}{AR_s}$$

and of the ratio

$$\frac{AR_{sd}}{SD} \frac{AR_d}{SS}$$

instead of

$$\frac{AR_{sd}}{AR_s}$$

were obtained from Group A and B (Table 1). The differences between the mean values obtained from Group A and that from Group B were statistically significant.

The correlation between

$$ds/dt$$

and

$$\frac{AR_s}{SB}$$

defined in Fig 2 can be seen in Fig 5.

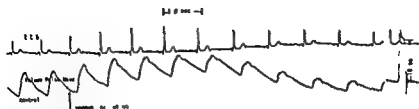


Fig. 6 Demonstration of waveforms obtained with venous occlusion. The venous occlusion cuff was applied to the arm just above the elbow crease and was inflated to a pressure less than diastolic.

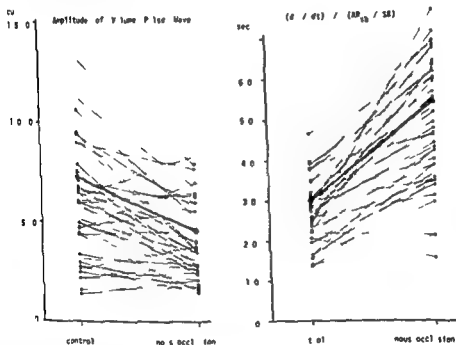


Fig. 7 Changes in the ratio $(d/dt)/(AR_v/SB)$ and in the amplitude of the pulse wave observed in Group A due to venous occlusion. Mean values of the ratio were elevated from 29.9 sec^{-1} to 53.8 sec^{-1} while those of the amplitude decreased from 7.3 mm^2 to 4.6 mm^2 .

considered as a constant during the single pulse cycle started at S . Equation (7) seems to represent clearly the displacement deflection of the volume pulse but it has an approximation involved in the application of Equations (4) and (5) to the digit that has non Hookean blood vessel with inertia and non ideal fluid flow.

According to the representation of the volume values reported by Burch²² five thousand mm^3 of finger tip the pulse wave of which was 7 mm^3 contained 800 mm^3 of blood. If the volume pulsation comes from venules and capillaries²³ the average velocities of the blood flow and pulse wave are about 2 cm per second and 200 cm per

second respectively.²³ The size of the pulse wave calculated by Equation (7) is 8 mm^3 ($\kappa = 1$) while reported size was 7 mm^3 . The calculated value is not very different from the actual value.

Let us try to interpret the meanings of the areas and ratios according to an application of Equation (7) to them. The areas can be defined as

$$AR_{ad} = \int_a^b \Delta r \cdot d = \frac{1}{c} \int_a^b \bar{U} \cdot dr \quad (8)$$

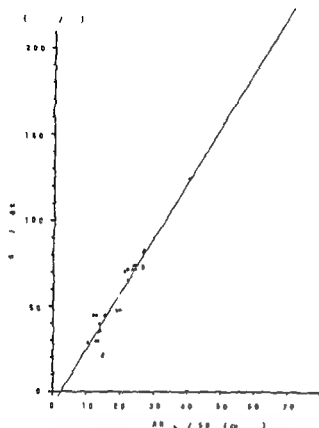


Fig 5 Correlation between (dV/dt) (Y axis) and AR_d/SB (X axis) in 93 pulse waves obtained from normal subjects $r = 0.69$ $P < 0.00005$ regression line $y = 3.3x - 9$

might be involved in the pulsatile change of the blood flow. That is, the higher amplitude of the volume pulse should be due to the increased volume flow. However there is no report that the blood flow of the digit is increased by raising the arm. On the contrary there are many reports that the blood flow was reduced when the digit was raised.^{17,18} Lack of correlation between blood flow and pulsation volume is often observed.¹⁹ It should be noticed that the pulsation volume does not indicate blood flow itself. One more disadvantage of Equation (1) is that it is very difficult to interpret the meanings of the areas and ratios

$$AR_d, AR, AR_d, \frac{AR_d SS}{AR_d SD}, \frac{AR_d DS'}{AR_d SD}$$

and

$$\frac{dV/dt}{AR_d}, \frac{dV/dt}{SB}$$

Of course it is convenient to have a theory which can explain everything concerning volume pulse. One theoretical proposal for

this purpose will be presented in the following section.

Let us consider a single straight cylindrical blood vessel instead of a network of small vessels in the digit studied. The resting radius of the cylindrical vessel, its longitudinal length, and its radial dilatation are denoted by R , L and ξ , respectively.

The resting volume of the vessel (V) is given by

$$V = \pi R^2 L \quad (2)$$

The volume increment of the vessel following radial dilatation due to pulsatile flow is given by

$$\Delta V = \pi (2 \xi R + \xi^2) L \quad (3)$$

According to Womersley²⁰ the radial dilatation is related to the velocity of flow, thus

$$\frac{2 \xi}{R} = \frac{\bar{U}}{c} \quad (4)^*$$

where \bar{U} is the average velocity of the blood flow and c is the wave velocity. From Equations (3) and (4) if $\bar{U}/4c < 1$ then the volume change of the vessel is given by

$$\Delta V = \frac{\pi R^2 L \bar{U}}{c} \quad (5)$$

It is obvious that the volume change of the digit (Δv) depends on the volume change of the blood vessel (ΔV) in it. The volume change of the digit—the displacement deflection of the volume pulse wave—is given by

$$\Delta v = \kappa \Delta V \quad (6)$$

where κ is a constant which relates to the elasticity of the tissue surrounding the vessel.

From Equations (2), (5) and (6) the displacement deflection of the volume pulse at a given moment (τ) during a single pulse cycle started at S can be written by

$$\Delta v = \frac{\pi L \bar{U}}{c} \quad (7)$$

The resting volume of the vessel is con-

Derivation of the equation is based on the ideal thin walled Hookean tube neglecting the inertia of the wall. It is also assumed that the radius of the tube in equilibrium under internal pressure is equal to its radius under zero transwall pressure.^{21,22}

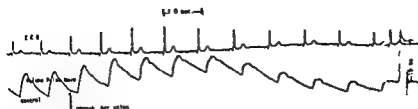


Fig. 6 Demonstration of waveforms obtained with venous occlusion. The venous occlusion cuff was applied to the arm just above the elbow crease and was inflated to a pressure less than diastolic.

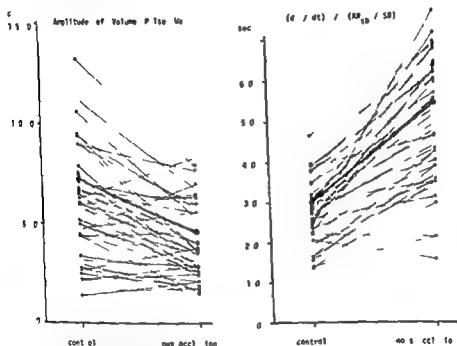


Fig. 7 Changes in the ratio $(d\epsilon/dt)/(AR_0/SB)$ and in the amplitude of the pulse waves observed in Group 1 due to venous occlusion. Mean values of the ratio were elevated from 29.9 sec^{-1} to 53.8 sec^{-1} while those of the amplitude decreased from 7.3 mm^3 to 4.6 mm^3 .

considered as a constant during the single pulse cycle started at S . Equation (7) seems to represent clearly the displacement deflection of the volume pulse but it has approximation involved in the application of Equations (4) and (5) to the digit that has non Hookean blood vessel with inertia and non ideal fluid flow.

According to the representation of the volume values reported by Burch²² five thousand mm^3 of finger tip the pulse wave of which was 7 mm^3 contained 800 mm^3 of blood. If the volume pulsation comes from venules and capillaries¹⁹ the average velocities of the blood flow and pulse wave are about 2 cm per second and 200 cm per

second respectively.²³ The size of the pulse wave calculated by Equation (7) is 8 mm^3 ($\kappa = 1$) while reported size was 7 mm^3 . The calculated value is not very different from the actual value.

Let us try to interpret the meanings of the areas and ratios according to an application of Equation (7) to them. The areas can be defined as

$$AR_{ad} = \int_a^D \Delta \sigma \, dr = \frac{\kappa}{c} \int_a^D \bar{U} \, d \quad (8)$$

Table 1 Mean values of the ratios obtained from two groups

| | Group A | Group B | |
|---------------------------|----------------|----------------------------|----------------------------|
| | Normal control | Cases with cardiac disease | Cases with cardiac failure |
| $\frac{AR_d DS}{AR_d SD}$ | 2.05 ± 0.43* | 1.83 ± 0.28* | 1.61 ± 0.21† |
| $\frac{AR_d SS}{AR_d SD}$ | 1.49 ± 0.15‡§ | 1.45 ± 0.16‡ | 1.29 ± 0.10‡ |

*Difference statistically significant to $P < 0.05$ †Difference statistically significant to $P < 0.005$ ‡Difference statistically significant to $P < 0.0001$

§Difference not statistically significant

Table 11 Mean values of $(dv/dt) / (AR_{ab}/SB)$ obtained from two groups

| | Group A | Group B | |
|------------------|----------------|----------------------------|----------------------------|
| | Normal control | Cases with cardiac disease | Cases with cardiac failure |
| Control | 29.9 ± 5.1**‡ | 31.9 ± 9.5*§ | 34.5 ± 9.1† |
| Venous occlusion | 53.8 ± 19.7‡ | 37.9 ± 11.0‡ | 37.6 ± 13.0 |

*Difference not statistically significant

†Difference not statistically significant

‡Difference statistically significant to $P < 0.00005$ (high significance)

§Difference not statistically significant

||Difference not statistically significant

and

$$AR_d = \int_D^S \Delta r \, d\tau = \frac{\kappa V}{c} \int_D^S \bar{U} \, d\tau \quad (10)$$

If we define the mean values of average velocity during the period of SD , SS' and DS' , (\bar{U}_d , \bar{U}_{ss} , and \bar{U}_{ds}) as

$$\bar{U}_d = \frac{1}{SD} \int_D^S \bar{U} \, d\tau$$

$$\bar{U}_{ss} = \frac{1}{SS'} \int_S^{SS'} \bar{U} \, d\tau$$

and

$$\bar{U}_{ds} = \frac{1}{DS} \int_D^{DS} \bar{U} \, d\tau$$

Equations (8), (9) and (10) can be reduced to

$$AR_{sd} = \frac{\kappa V \bar{U}_{sd} SD}{c} \quad (11)$$

$$AR = \frac{\kappa V \bar{U}_{ss} SS}{c} \quad (12)$$

and

$$AR_d = \frac{\kappa V \bar{U}_d DS}{c} \quad (13)$$

respectively

From Equations (11), (12) and (13) (13) the ratios are given by

$$\frac{AR_{sd}}{AR} = \frac{L_{sd}}{L} \quad (14)$$

and

$$\frac{AR_{sd}}{AR_s} = \frac{L_{sd}}{L_s} \quad (15)$$

where L_s and L_{sd} are the mean displacements of the blood during SD , SS , and DS respectively. It is obvious that they closely correlate with one another. One may also easily obtain the ratios of velocities of blood flow corresponding to

$$\frac{SD}{SS}$$

and

$$\frac{SD}{DS}$$

as follows

$$\frac{AR_{sd} SS}{AR SD} = \frac{\bar{U}_{sd}}{\bar{U}} \quad (16)$$

and

$$\frac{AR_{sd} DS}{AR SD} = \frac{\bar{U}_{sd}}{\bar{U}_s} \quad (17)$$

The difference between the mean values of the ratios of the normal persons and those with cardiac disease or cardiac failure can be explained as the difference between the mean values of average velocities during SD and those during DS corresponding to each group.

Next application of Equation (7) is to AR_s which is the area under the volume pulse over SB interval. That is

$$AR_s = \int_a^b \Delta v \, dt = \frac{\pi V}{t} \int_a^b \bar{U} \, dt \quad (18)$$

If the mean value of the average velocity of flow during SB (\bar{U}_s) is defined as

$$\bar{U}_s = \frac{1}{SB} \int_a^b \bar{U} \, dt$$

Equation (18) can be reduced to

$$AR_s = \frac{\pi \bar{U}_s SB}{c} \quad (19)$$

The amplitude of the volume pulse at a point p that was shown in Fig. 2 can be expressed as

$$\Delta v_p = \frac{\pi V \bar{U}_p}{c} \quad (20)$$

The maximum rate of rise in the pulse wave at the point p is given by

$$\lim_{\Delta t \rightarrow 0} \frac{\Delta v}{\Delta t} = \frac{dv}{dt} = \frac{\pi V}{c} \frac{d\bar{U}_p}{dt} \quad (21)$$

Thus the acceleration of the blood flow is due to the influences of some unbalanced force (F) on the blood mass (m) in the cylindrical vessel. The force acting on the blood mass at the moment p can be described as

$$F = \frac{d(m \bar{U}_p)}{dt} = \frac{mc}{t} \frac{dv}{dt} \quad (22)$$

If force (F) is expressed by pressure (P)

$$P = \frac{cm}{t} \frac{dv}{dt} \quad (23)$$

where A is the total area on which the force is acting.

From Equations (19) and (23) the ratio of

$$\frac{dv}{dt}$$

to

$$\frac{AR_s}{SB}$$

is given by

$$\frac{dv/dt}{AR_s/SB} = \frac{A}{m} \frac{P}{\bar{U}_s} \quad (24)$$

According to the Poiseuille law governing laminar flow in a cylindrical tube, pressure drop (P) is expressed as

$$P = \frac{8 \mu L \bar{U}}{R^2} \quad (25)$$

where μ is the viscosity, L the length of the vessel, \bar{U} the mean velocity of flow, and R the radius of the vessel. If Poiseuille's formula is applicable to the steady flow component—the mean flow—of the pulsatile arterial flow, Equation (24) can be rewritten as

$$\frac{dv/dt}{AR_s/SB} = \frac{K}{m} \frac{A}{m} \quad (26)$$

where K is the peripheral resistance.

The reason for the constancy of the ratio

$$\frac{dv/dt}{AR_s/SB}$$

is due to the constancy of the peripheral resistance (Fig 5). Highly significant differences between the mean values of ratio observed in normal control and those changed by venous occlusion can be explained by the increase of the peripheral resistance that is probably due to a drop in the mean velocity of the blood flow (Fig 7). In Group B, values of the ratio were not significantly changed by venous occlusion because of the underlying high peripheral resistance in the cases of persons with cardiac disease or cardiac failure (Table II).

The effect of venous occlusion on the amplitude of the volume pulse can be explained by a decrease of flow velocity and an increase of wave velocity. It is clear that the decrease in the flow velocity is caused by a decreased pressure gradient between the mean pressure of the arteriole and that of the venule and the increased peripheral resistance as shown in the elevation of the ratio

$$\frac{dv/dt}{AR_v/SB}$$

(Fig 7). The wave velocity is expressed by Bramwell and Hill¹⁶ as follows

$$c = k \sqrt{V \frac{dp}{dv}}$$

where k is a constant and V is the initial arterial volume per unit length. The increase in the wave velocity can be clearly explained by the increased bulk modulus

$$V \frac{dp}{dv}$$

in the equation due to venous congestion.

When the digit is raised above the heart the amplitude of the pulse wave should become larger because of the decreased velocity of the pulse wave and a slightly increased velocity in the flow. It is clear that the velocity of the pulse wave becomes slower because of a reduction in the bulk modulus. As reported by Holling and Verel,¹⁷ the fall in systolic and diastolic arterial pressure that accompanies an elevation of the arm corresponds to the change in hydrostatic pressure. The pulse pressure does not significantly alter as the arm is lifted. When the arm is lifted, the venous pressure falls slightly but not below 1 to

—3 mm Hg, though the brachial venous pressure is 2 to 5 mm Hg at heart level. These findings indicate that the pressure gradient between the mean pressure of the arteriole and that of the venule remains constant or slightly increases with an elevation of the arm. The ratio

$$\frac{(dv/dt)}{(AR_v/SB)}$$

indicating peripheral resistance does not alter by an elevation of the arm (Fig 5). Therefore it might be considered that the average velocity of the flow is almost equal to the control or is slightly increased when the arm is lifted.

The deviation of the base line of the pulse wave seen in the venous occlusion method can be explained by the increment of the resting volume (V) in Equation (7). The principle of the method might be based on the possibility that the increment of the resting volume caused by arterial inflow under no venous return must be proportional to the degree of deviation of the base line. If the ratio of flow velocity to wave velocity is not significantly altered by the venous occlusion it is possible to measure blood flow by the method.

Summary

One simple equation is proposed for a representation of the volume pulse wave

$$\Delta v = \frac{\pi V \bar{U}}{c}$$

where Δv is the volume change of the digit studied—i.e. the amplitude of the pulse wave at a given moment τ ; π constant V the resting volume of the blood vessel \bar{U} the average velocity of the blood flow at the given moment and c the wave velocity.

$$\frac{AR_s}{SD} \quad \frac{AR_d}{DS}$$

and

$$\frac{AR}{SS}$$

are closely correlated with each other. The constancy of the ratios

$$\frac{AR_s}{SD} / \frac{AR_d}{DS} \quad \frac{AR_s}{SD} / \frac{AR}{SS}$$

can be explained as follows

$$\frac{AR_{ss}}{AR_{SD}} = \frac{\bar{U}_{ss}}{\bar{U}_{SD}}$$

$$\frac{AR_{DS}}{AR_{SD}} = \frac{\bar{U}_{DS}}{\bar{U}_{SD}}$$

where \bar{U}_{ss} , \bar{U}_{DS} and \bar{U}_{SD} are the mean values of the average velocity of the flow during SD , DS and SS respectively

The ratio of the rate of rise of the pulse wave

$$dv/dt$$

to

$$\frac{AR_{ss}}{SB}$$

is also constant under normal conditions AR_{ss} signifies the area under the pulse wave during SB . The ratio can be expressed as

$$\frac{dv/dt}{AR_{ss}/SB} = \frac{K}{m} \cdot A$$

where K is the peripheral resistance, A the total area on which the force is acting and m the blood mass on which the force is acting

To obtain the values of \bar{U} , c and K having clinical significance through the analysis of the volume pulse further advance of the method should be expected

We are very grateful to Dr Ichiro Matsubara for his valuable advice

REFERENCES

1. Quoted from Allen E V, Barker N W and Hines E A Jr. *Peripheral vascular disease*. Philadelphia 1946. W B Saunders Company p 130
2. Grant R T and Bruce Pearson R S. The blood circulation in the human limb: observations on the differences between the proximal and distal parts and remarks on the regulation of body temperature. *Clin Sci* 3:119 1937 1938
3. Hertzman A H. The blood supply of various skin areas as estimated by the photoelectric plethysmograph. *Am J Physiol* 124:328 1938
4. Abramson D I, Zazetla H and Marrus J. Plethysmographic studies of peripheral blood flow in man: I. Criteria for obtaining accurate plethysmographic data. *AM HEART J* 17 1934 1939
5. Burch G E, Cohn A E and Neumann C. A study by quantitative methods of the spontaneous variations in volume of the finger tip, toe tip and postero-superior portion of the pinna of resting normal white adults. *Am J Physiol* 136:133 1947

6. Landowne M and Katz L N. A critique of the plethysmographic method of measuring blood flow in the extremities of man. *AM HEART J* 23:644 1942
7. Goetz H H. The rate and control of the blood flow through the skin of the lower extremities. *AM HEART J* 31:146 1946
8. Nyboer J. Electrical impedance plethysmography. A physical and physiologic approach to peripheral vascular study. *Circulation* 2:811 1950
9. Winor T. Clinical Plethysmography Part I. An improved direct writing plethysmograph. *Angiology* 4:134 1953. Part II. Plethysmographic procedures of clinical importance. *Angiology* 4:149 164 1953
10. Clarke R S J and Mellon R F. Venous collection in forearm and hand measured by the strain gauge and volume plethysmograph. *Clin Sci* 16:103 1957
11. Winsor T. The segmental plethysmograph. A description of the instrument. *Angiology* 8:57 1957
12. Brodie T G and Russell A E. On the determination of the rate of blood flow through an organ. *J Physiol* 32:17 1905
13. Greenfield A D M, Whitney R J and Mowbray J F. Methods for the investigation of peripheral blood flow. *Br Med Bull* 19 101 1963
14. Ardill B L, Bhatnagar V M, Fentem P H and Greenfield A D M. Clinical use of venous occlusion plethysmography. *Scand J Clin Lab Invest (Suppl 99)* 95 1967
15. Nyboer J. *Electrical impedance plethysmography*. Springfield Ill 1970. Charles C Thomas Publisher p 145
16. Burch G E. Method for recording simultaneously the time course of digital rate and of digital volume of inflow, outflow and the difference between inflow and outflow during a single pulse cycle in man. *J Appl Physiol* 7:99 1954
17. Holling H H and Verel H. Circulation in the elevated forearm. *Clin Sci* 16:197 1957
18. Gaskell P and Burton A. Local postural vasomotor reflexes arising from the limb veins. *Circ Res* 1:27 1953
19. Mune O. Pulse morphological plethysmography of the forefoot in arteriosclerosis obliterans. *Scand J Clin Lab Invest (Suppl 99)* 147 1967
20. Womersley J R. Oscillatory motion of a viscous liquid in a thin walled elastic tube I. The linear approximation for long waves. *Philosophical Magazine* 46:199 1955
21. Womersley J R. Oscillatory flow in arteries: the constrained elastic tube as a model of arterial flow and pulse transmission. *Phys Med Biol* 2:178 187 1957
22. Burch G E. A new sensitive portable plethysmograph. *AM HEART J* 33:48 1947
23. Berne R M and Levy M N. *Cardiovascular*

- physiology St Louis 1967, The C V Mosby Company p 3
- 24 Karremann G. Some contributions to the mathematical biology of blood circulation. Reflections of pressure wave in the arterial system Bull Math Biophys 14:327, 1952
- 25 McDonald D A. Blood flow in arteries Baltimore 1960 The Williams & Wilkins Company p 178
- 26 Bramwell C and Hill A V. The velocity of the pulse wave in man Proc R Soc Lond (Biol) 93:298 1922

Infection of an avulsed papillary muscle tip simulating bacterial endocarditis

Terry K. Satterwhite MD

Zell A. McGee MD*

William Schaffner MD

Gottlieb C. Friesinger MD

Mona Mishu MD

Robert D. Collins MD

Nashville Tenn

Rupture of a papillary muscle usually initiates a series of events that terminates within hours or days in cardiac decompensation and death.^{1,2} However, a few patients have developed only moderate mitral insufficiency and lived for months with a ruptured papillary muscle.³ This report describes a patient who had a ruptured papillary muscle for an indeterminate length of time, this condition being called to medical attention by the presence of gram negative bacteremia which persisted despite appropriate antimicrobial therapy. Clinically this was thought to represent bacterial endocarditis with ruptured chordae tendineae. At postmortem examination the patient's illness was found to be caused by persistent infection of an avulsed papillary muscle tip.

Case report

The patient was a 74 year old man with diabetes. Six weeks prior to admission he noted the onset of

anorexia, nausea, postprandial fullness and weight loss. Three weeks later he began having diaphoresis, chills, and fever to 106° F. A white blood cell count of 10,000 per mm³ and a hemoglobin of 13 Gm per 100 ml were observed at his local hospital. Five days of intravenous tetracycline failed to alter his course and he was admitted to the Vanderbilt University Hospital.

Mild dyspnea on exertion had been present for several months but the patient denied orthopnea, paroxysmal nocturnal dyspnea, or chest discomfort. There was no history of rheumatic fever or myocardial infarction. A soft systolic murmur at the apex was present a year prior to admission.

The diagnosis of diabetes mellitus had been made 10 years previously and he had taken insulin for six years. He was receiving no other medicine at the time of admission. A left below the knee amputation had been done 15 months earlier because of vascular insufficiency. Urinary frequency and hesitancy had been present for several years.

Physical findings included a supine blood pressure of 104/54 mm Hg, a regular pulse of 92 per minute, and a rectal temperature of 101° F. Splinter hemorrhages, conjunctival petechiae, and splenomegaly were noted. Examination of the chest revealed no abnormalities. There was no jugular venous distention at 45 degrees. The heart was not enlarged.

From the Division of Infectious Disease and the Division of Cardiology, Department of Medicine and the Department of Pathology, Vanderbilt University School of Medicine, Nashville, Tenn.
Supported in part by Grant AI 03092 and AI 03323 from the National Institute of Allergy and Infectious Diseases and a grant from the Lilly Laboratory for Clinical Research and Education.
Received for publication August 12, 1972.
Reprint requests: Zell A. McGee, MD, Division of Infectious Diseases, Vanderbilt University Medical Center, Nashville, Tenn. 37232.
Dr. McGee's present address: Department of Infectious Diseases, Developmental and Clinical Pharmacology, National Institute of Allergy and Infectious Diseases.

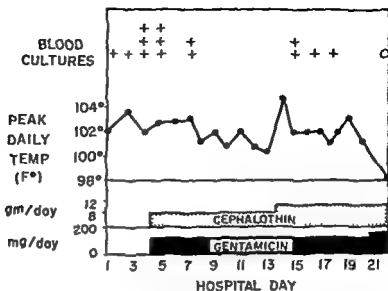


Fig 1 Schematic representation of clinical course

The first heart sound and a systolic click were palpable. The click occurred early in systole and was followed by a harsh III/VI murmur that lasted throughout the rest of systole. The murmur was heard along the left sternal border and at the cardiac apex with radiation to the axilla. There were no bruits over the stump of the left lower extremity. Right costovertebral angle tenderness was present. The prostate was firm and enlarged.

Initial laboratory studies revealed a white blood cell count of 10 100 per mm^3 with 89 per cent segmented forms. Some of the neutrophils contained toxic granulations. The hematocrit was 36 per cent, platelet count 333 000 per mm^3 , and the latex fixation test for rheumatoid factor was negative. The blood urea nitrogen was 23 mg per 100 ml and glucose 370 mg per 100 ml. Examination of the urine showed 4+ sugar without ketone. There was microscopic hematuria.

Two blood cultures drawn during the first 24 hours after admission were positive for *Klebsiella pneumoniae* (Fig 1) and the admission urine culture also yielded *Klebsiella pneumoniae* in a concentration greater than 100 000 organisms per ml with identical sensitivities to the organism in the blood.

The clinical impressions were that the patient had *Klebsiella* endocarditis with ruptured chordae tendineae resulting in mitral insufficiency. Therapy was instituted with intramuscular gentamicin 5 mg per kilogram of body weight per day and intravenous cephalothin 8 gm per day. Some of the laboratory and clinical features of his hospital course are presented in Fig 1. The minimal bactericidal concentrations of cephalothin and gentamicin for the *Klebsiella* isolated from the patient's blood were 62 μg per milliliter and 0.39 μg per milliliter respectively. His serum was bactericidal for the *Klebsiella* when diluted 1:8 on the fourth day of therapy and 1:16 on the sixth day of therapy. This indicated that serum levels of antimicrobial activity usually considered adequate in treating endocarditis were being achieved. Nevertheless, the patient remained febrile and blood cultures were repeatedly positive for

Klebsiella. Subsequent urine cultures were negative.

Daily examination of the patient's abdominal films, two intravenous pyelograms, a barium enema, gall bladder series, thoracolumbar spine films, and a liver scan failed to reveal a source of bacteremia other than the heart. A progressive deterioration in renal function ensued with the serum urea nitrogen and creatinine reaching 72 mg per 100 ml and 7.8 mg per 100 ml, respectively. Because of the severity of the underlying medical problems, mitral valve replacement was deferred and antibiotic therapy was continued.

The admission ECG showed normal sinus rhythm with low voltage in the limb leads and non-specific ST-T wave changes. On the twentieth hospital day, he was noted for the first time to have an irregular pulse. An ECG at this time revealed supraventricular premature contractions and short bursts of supra-ventricular tachycardia with aberrant ventricular conduction. Digitalis was administered cautiously. On the next day, he became hypotensive and died.

At necropsy, the heart weighed 300 gm. The right and left atria were moderately dilated but the other chambers were not enlarged. The aortic pulmonary and tricuspid valves were not remarkable. The mitral valve ring measured 8.5 cm and the valve appeared normal with no gross evidence of recent or remote endocarditis. All mitral leaflet chordae tendineae were attached and were not thickened or inflamed. The aortic leaflet of the mitral valve was remarkable in that the most central chordae were not fixed to the ventricular surface due to an avulsion of part of the papillary muscle (Fig 2).

The avulsed piece of papillary muscle measured 11 by 7 by 6 mm, was yellowish white, and was attached to the valve by normal appearing chordae tendineae. The previous point of attachment of this portion of papillary muscle was not readily apparent on gross examination. Both anterolateral and posteromedial papillary muscles had smooth surfaces and showed minimal focal subendocardial fibrosis. However, microscopic examination of the anterolateral papillary muscle revealed remote myocardial

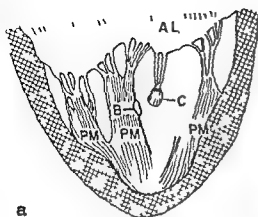


Fig 2 a through c a Schematic illustration and photograph of gross specimen of patient's heart. (a) anterior section through the left ventricle illustrating the ruptured papillary muscle tip (C) attached to the aortic leaflet of the mitral valve (AL) by its chordae tendineae. B indicates the base of the papillary muscle at the site of rupture. Papillary muscles are labeled PM. b Photomicrograph showing the base of the papillary muscle at the site of rupture. There are no bacteria and there is no evidence of inflammatory reaction (Hematoxylin and eosin. Original magnification $\times 435$). c Photomicrograph demonstrating gram-negative bacilli in the center of the avulsed tip and the absence of inflammatory infiltrate (Gram's stain. Original magnification $\times 2,700$).

necrosis, calcification and mild chronic inflammation, presumably at the site of avulsion of the papillary muscle fragment (Fig 2). Search of multiple sections of the anterolateral papillary muscle base showed no abscesses and no bacteria were seen on gram-stained sections (Fig 2b). In contrast the avulsed papillary tip was teeming with gram-negative bacilli (Fig 2c). No inflammatory cells were seen among the bacteria nor was any viable muscle found. The tissue of the avulsed tip was eosinophilic and had the appearance of autolyzed cardiac muscle. Despite the lack of support due to the avulsed papillary muscle tip, the aortic leaflet of the mitral valve was anchored by other chordae tendineae from both the anterior and posterior papillary muscles.

The coronary arteries showed calcification and marked atherosclerosis. Although narrowed, no point of obstruction or thrombosis was found. Multiple sections of the left ventricular myocardium failed to reveal recent or remote infarction.

Each kidney weighed 200 Gm with multiple small scars over the surface. Microscopic section of

the kidneys showed arterial and arteriolar nephrosclerosis. There was chronic inflammation present in the cortex and medulla of both kidneys. The prostate was enlarged and showed glandular hyperplasia and microabscesses.

Comments

The patient was admitted to the hospital with a history very suggestive of bacterial endocarditis. The physical findings of a heart murmur, fever, petechiae, enlarged spleen, anemia and splinter hemorrhages were especially convincing. This impression was strengthened by multiple positive blood cultures and a urine culture positive for *Klebsiella*. The infection, however, was persistent in spite of appropriate treatment and only at necropsy was its location found to be the avulsed papillary muscle tip. In

attempting to correlate the necropsy findings with his clinical course several aspects of the illness deserve comment. The etiology of the disease of the mitral apparatus is especially perplexing.

The etiologies, pathology and pathophysiology of mitral regurgitation have been a topic of much interest in recent years. Rheumatic mitral regurgitation with pathological involvement of the mitral leaflets has become proportionately much less frequent as the spectrum of mitral regurgitation has become better defined. Any portion of the mitral valve apparatus or its attachments may become diseased in a manner to create regurgitation. Physical findings and the clinical course may be characteristic for a given etiology.

Papillary muscle dysfunction is an extremely common etiology for mitral regurgitation, ischemic heart disease being the entity usually responsible for this variety of mitral regurgitation. The murmur of papillary muscle dysfunction is non holosystolic and usually relatively localized to the apical area. Ordinarily few symptoms are present and endocarditis is not a complication. Papillary muscle rupture occurs most often in association with acute myocardial infarction and is characteristically associated with a pan systolic murmur of great intensity. It is customary for profound left ventricular failure to occur at the time of rupture. Elongation of the chordae tendineae can account for mitral regurgitation which is usually associated with characteristic physical findings including a systolic click and a crescendo-decrescendo systolic murmur. These patients ordinarily do not have symptoms. Trauma is a very rare cause for isolated chordae rupture.

Our patient does not fit characteristically into any of the categories of mitral regurgitation described above. The documentation of a systolic murmur prior to the onset of the illness suggests that the papillary muscle ruptured before the infective process developed. This is in contrast to an instance of myocardial abscess reported by Hackel and Kaufman.³ In their patient, a new mitral murmur appeared and the patient rapidly deteriorated. Autopsy demonstrated that the papillary muscle was detached from the ventricle. Microscopic sections of

both the ventricle and papillary muscle revealed abscess formation at the edges of the separation with gram negative bacteria being present.

The separated papillary muscle tip of our patient was found to have undergone complete necrosis and was without inflammatory reaction. Many gram negative bacilli were present in this avulsed papillary tip. In contrast and unlike the patient reported by Hackel and Kaufman,³ neither necrosis, inflammatory reaction, nor bacteria were present in the base of the papillary muscle at the site of separation of the tip. Hence it seems unlikely that an abscess with subsequent rupture of the papillary muscle is the explanation in this case. It is probable that the papillary muscle rupture antedated the infection. This would explain why his illness presented principally as an infectious disease and not importantly as myocardial dysfunction.

Two findings may explain the mild hemodynamic effects of the ruptured papillary muscle. (1) the aortic leaflet was anchored by chordae tendineae from the remainder of the anterolateral as well as the posteromedial papillary muscles and more importantly (2) the fragment that ruptured from the main body of the papillary muscle was small so that only 2 of the 18 chordae that anchored the mitral valve apparatus were lost. The resulting dysfunction of the mitral valve was apparently minimal and was indicated only by the systolic click and the murmur of mitral insufficiency rather than by congestive heart failure.

A postulated sequence of events proceeds as follows: the small papillary muscle tip ruptured but caused only minimal hemodynamic disturbance. The patient's chronic *Klebsiella* urinary tract infection produced a transient *Klebsiella* bacteremia which infected the free floating tip. Because of its avascular nature the tip acted as an infected foreign body analogous to a prosthetic heart valve. In such avascular tissues adequate tissue levels of antibiotics are frequently not obtained and phagocytes are unable to gain access to the lesion. Thus sustained bacteremia continued despite the presence of what are usually considered to be adequate serum bactericidal levels of antibiotics to which the organism was sensitive. Eradication of infections in such areas

cular structures is extremely difficult and usually requires antibiotic therapy plus surgical removal of the intravascular foreign body.⁴

To our knowledge this is the first report of a ruptured papillary muscle presenting clinically as bacterial endocarditis and sustained bacteremia. Our experience has been similar to that of McHenry and associates⁷ in that sustained bacteremia despite appropriate antimicrobial therapy has usually been associated with undrained abscesses, infection of foreign bodies such as prosthetic heart valves contaminated in travenous fluids and lines, or severe burns. When a sustained bacteremia despite appropriate antimicrobial therapy occurs in association with mitral insufficiency, serious consideration should be given to the possibility of an infection of an avascular portion of the mitral apparatus.

We thank Dr Howard E. Rosen for permitting us to report this patient.

REFERENCES

- 1 Sanders R J, Neuburger K T and Ravin A. Rupture of papillary muscles. Occurrence of rupture of the posterior muscle in posterior myocardial infarction. *Dis Chest* 31:316 1957
- 2 Austen W G, Sokol D M, DeSanctis R W and Sanders C A. Surgical treatment of papillary muscle rupture complicating myocardial infarction. *N Engl J Med* 278:1137 1968
- 3 DeBusk R F and Harrison D C. The clinical spectrum of papillary muscle disease. *N Engl J Med* 281:1458 1969
- 4 Breneman G M and Drake F H. Ruptured papillary muscle following myocardial infarction with long survival. *Circulation* 28:867 1962
- 5 Hackel D H and Kaufman N. Papillary muscle rupture due to a myocardial abscess. *Ann Intern Med* 38:824 1953
- 6 Killen D A, Collins H A, Koenig M G and Goodman J S. Prosthetic cardiac valves and bacterial endocarditis. *Ann Thorac Surg* 9:238 1960
- 7 McHenry M C, Gavan T L, Hawk W A and Vanomem C A. Gram negative bacteremia of long duration. Eleventh Interscience Conference on antimicrobial Agents and Chemotherapy (Abstr) 12 1971

Left-to-right shunt at atrial level after rupture of papillary muscle from acute myocardial infarction

Michael R Nagel, MD*

James A Ronan, Jr, MD**

William C Roberts MD***

Washington D C

A systolic murmur may develop during or after acute myocardial infarction from ischemia, necrosis, fibrosis, or rupture of a papillary muscle or from rupture of the ventricular septum. Differentiation of mitral regurgitation from ventricular septal rupture in this setting may be difficult but the distinction is often made by the detection of a high oxygen content of blood in the pulmonary artery strongly suggesting the presence of a ventricular septal defect. Rarely, mitral regurgitation produces retrograde flow through the pulmonary capillary bed causing elevation of the pulmonary arterial oxygen content.¹ We recently studied a patient who, during acute myocardial infarction developed increased blood oxygen content in the right side of the heart due to papillary muscle rupture and a newly acquired left to right shunt at atrial level. Hemodynamic data indicated that

the shunt was located at the atrial level, and necropsy showed that the defect was due to stretching of the fossa ovalis region by left atrial dilatation after the acute development of mitral regurgitation. Pertinent clinical and anatomic features of this patient are summarized in this report.

Case report

W A (021 39 50) a 65 year old man who died August 13 1970 was first seen at Georgetown University Hospital on August 4 1970. An acute diaphragmatic wall myocardial infarction had occurred 75 days earlier when he had been admitted to another hospital. Six days after the onset of infarction he developed a loud holosystolic apical murmur and severe congestive cardiac failure. Despite bed rest digitalis diuretics and salt restriction during the next 69 days he remained in chronic heart failure and became cachectic. The electrocardiograms (ECGs) taken by his private physician in previous years specifically showed none of the ECG features of atrial septal defect and his previous chest roentgenograms were normal.

From the Department of Medicine Division of Cardiology Georgetown University School of Medicine Washington D C

This work was supported by grants from the U S Public Health Service the Benjamin May Memorial Fund and the Special Cardiac Fund

Received for publication Aug 14 1972

Reprint requests to Dr James A Ronan Jr Division of Cardiology Georgetown University 1100 3800 Reservoir Rd NW Washington D C 20007

*Fellow in Cardiology Division of Cardiology Georgetown University Hospital Washington D C Present address 15955 Samaritan Dr San Jose Calif 94125

**Associate Professor of Medicine Division of Cardiology Georgetown University School of Medicine Washington D C

***Chief Section of Pathology National Heart and Lung Institute National Institutes of Health Bethesda Md 20014 and Clinical Associate Professor of Pathology and Medicine (Cardiology) Georgetown University Washington D C



Fig 1 Left ventricular cineangiogram in the left anterior oblique view. There is a herniation of the atrial septum (arrows) with contrast material crossing from the left to the right atrium. Radioopaque dye can be seen in the left ventricle, aorta, and left atrium. The visible catheter is in the right atrium and right ventricle, but the tip in the pulmonary artery is not visible. The left ventricular catheter cannot be seen.

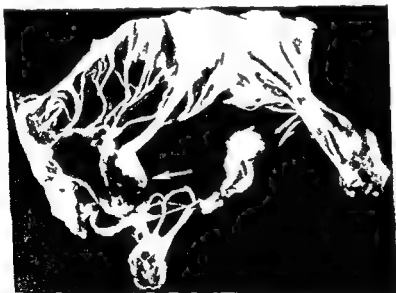


Fig 2 Surgical specimen of the mitral valve leaflets and supporting structures. That portion of the posteromedial papillary muscle which had ruptured is smooth and endothelialized (arrow).

Right heart catheterization at Georgetown University Hospital showed pulmonary hypertension and disproportionate elevation of the pulmonary arterial wedge V waves indicating mitral regurgitation (Table I). The hydroxyl inhalation test with the platinum tipped electrode in the right atrium yielded an appearance time of 3 seconds, documenting a left-to-right shunt at atrial level. Krypton 85 inhalation test with simultaneous blood sampling from pulmonary and left brachial arteries

(on two occasions) yielded a calculated pulmonary-to-systemic flow ratio in excess of 4:1. Left ventricular cineangiography disclosed moderately severe mitral regurgitation. A well-defined portion of the atrial septum bulged into the right atrium, and contrast material passed from the left atrium to the right atrium at that position (Fig 1).

At operation on August 13, 1970, portions of the posteromedial papillary muscle were found detached from its main trunk (Fig 2). The ruptured

Left-to-right shunt at atrial level after rupture of papillary muscle from acute myocardial infarction

Michael R. Nugent, M.D.*

James A. Henson, Jr., M.D.*

William L. Roberts, M.D. + Co.

Washington, D. C.

A syntactic movement may develop during an utterance, may not develop from the initial, pre-lexical, stage to, then to, an interpretation of a proposition, may be an interpretation of the semantic content, a different interpretation of an initial interpretation from semantic content, or may be a different way to state the same content, but the choice from the latter two by the choice of a single expression content of the latter in the preliminary action, strongly suggesting the presence of a semantic content choice. Finally, initial interpretation provides a way to state the content of the preliminary interpretation but a change of content of the preliminary interpretation content. We previously noted that a path of an utterance is the pre-lexical initial stage, developed to a semantic content in the right side of the first stage to a proposition interpretation and a way to express the content of the first stage at a higher level. Dynamically, this indicates that

the third was located at the trial level, and in response showed that the defect was due to a violation of the fair trial system by the trial magistrate after the war. The report of initial investigation did not establish any material features of the problem and was insufficient in this regard.

Cash report

W. A. (D) (1940), a 64 year old man who died August 14, 1970, was first seen at the Longview State Hospital in June, August 3, 1970. At the time of admission, wall space with his feet in the air 29 days a week when he had been admitted to see what his feet did after the onset of infection in the abdomen a hard knot, growth and pain and severe constipation, rather than the other two to 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 159, 160, 161, 162, 163, 164, 165, 166, 167, 168, 169, 170, 171, 172, 173, 174, 175, 176, 177, 178, 179, 180, 181, 182, 183, 184, 185, 186, 187, 188, 189, 190, 191, 192, 193, 194, 195, 196, 197, 198, 199, 200, 201, 202, 203, 204, 205, 206, 207, 208, 209, 210, 211, 212, 213, 214, 215, 216, 217, 218, 219, 220, 221, 222, 223, 224, 225, 226, 227, 228, 229, 230, 231, 232, 233, 234, 235, 236, 237, 238, 239, 240, 241, 242, 243, 244, 245, 246, 247, 248, 249, 250, 251, 252, 253, 254, 255, 256, 257, 258, 259, 260, 261, 262, 263, 264, 265, 266, 267, 268, 269, 270, 271, 272, 273, 274, 275, 276, 277, 278, 279, 280, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301, 302, 303, 304, 305, 306, 307, 308, 309, 310, 311, 312, 313, 314, 315, 316, 317, 318, 319, 320, 321, 322, 323, 324, 325, 326, 327, 328, 329, 330, 331, 332, 333, 334, 335, 336, 337, 338, 339, 340, 341, 342, 343, 344, 345, 346, 347, 348, 349, 350, 351, 352, 353, 354, 355, 356, 357, 358, 359, 360, 361, 362, 363, 364, 365, 366, 367, 368, 369, 370, 371, 372, 373, 374, 375, 376, 377, 378, 379, 380, 381, 382, 383, 384, 385, 386, 387, 388, 389, 390, 391, 392, 393, 394, 395, 396, 397, 398, 399, 400, 401, 402, 403, 404, 405, 406, 407, 408, 409, 410, 411, 412, 413, 414, 415, 416, 417, 418, 419, 420, 421, 422, 423, 424, 425, 426, 427, 428, 429, 430, 431, 432, 433, 434, 435, 436, 437, 438, 439, 440, 441, 442, 443, 444, 445, 446, 447, 448, 449, 450, 451, 452, 453, 454, 455, 456, 457, 458, 459, 460, 461, 462, 463, 464, 465, 466, 467, 468, 469, 470, 471, 472, 473, 474, 475, 476, 477, 478, 479, 480, 481, 482, 483, 484, 485, 486, 487, 488, 489, 490, 491, 492, 493, 494, 495, 496, 497, 498, 499, 500, 501, 502, 503, 504, 505, 506, 507, 508, 509, 510, 511, 512, 513, 514, 515, 516, 517, 518, 519, 520, 521, 522, 523, 524, 525, 526, 527, 528, 529, 530, 531, 532, 533, 534, 535, 536, 537, 538, 539, 540, 541, 542, 543, 544, 545, 546, 547, 548, 549, 550, 551, 552, 553, 554, 555, 556, 557, 558, 559, 560, 561, 562, 563, 564, 565, 566, 567, 568, 569, 570, 571, 572, 573, 574, 575, 576, 577, 578, 579, 580, 581, 582, 583, 584, 585, 586, 587, 588, 589, 590, 591, 592, 593, 594, 595, 596, 597, 598, 599, 600, 601, 602, 603, 604, 605, 606, 607, 608, 609, 610, 611, 612, 613, 614, 615, 616, 617, 618, 619, 620, 621, 622, 623, 624, 625, 626, 627, 628, 629, 630, 631, 632, 633, 634, 635, 636, 637, 638, 639, 640, 641, 642, 643, 644, 645, 646, 647, 648, 649, 650, 651, 652, 653, 654, 655, 656, 657, 658, 659, 660, 661, 662, 663, 664, 665, 666, 667, 668, 669, 670, 671, 672, 673, 674, 675, 676, 677, 678, 679, 680, 681, 682, 683, 684, 685, 686, 687, 688, 689, 690, 691, 692, 693, 694, 695, 696, 697, 698, 699, 700, 701, 702, 703, 704, 705, 706, 707, 708, 709, 710, 711, 712, 713, 714, 715, 716, 717, 718, 719, 720, 721, 722, 723, 724, 725, 726, 727, 728, 729, 730, 731, 732, 733, 734, 735, 736, 737, 738, 739, 740, 741, 742, 743, 744, 745, 746, 747, 748, 749, 750, 751, 752, 753, 754, 755, 756, 757, 758, 759, 760, 761, 762, 763, 764, 765, 766, 767, 768, 769, 770, 771, 772, 773, 774, 775, 776, 777, 778, 779, 780, 781, 782, 783, 784, 785, 786, 787, 788, 789, 790, 791, 792, 793, 794, 795, 796, 797, 798, 799, 800, 801, 802, 803, 804, 805, 806, 807, 808, 809, 810, 811, 812, 813, 814, 815, 816, 817, 818,

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61 62 63 64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 111 112 113 114 115 116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131 132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147 148 149 150 151 152 153 154 155 156 157 158 159 160 161 162 163 164 165 166 167 168 169 170 171 172 173 174 175 176 177 178 179 180 181 182 183 184 185 186 187 188 189 190 191 192 193 194 195 196 197 198 199 200 201 202 203 204 205 206 207 208 209 210 211 212 213 214 215 216 217 218 219 220 221 222 223 224 225 226 227 228 229 230 231 232 233 234 235 236 237 238 239 240 241 242 243 244 245 246 247 248 249 250 251 252 253 254 255 256 257 258 259 260 261 262 263 264 265 266 267 268 269 270 271 272 273 274 275 276 277 278 279 280 281 282 283 284 285 286 287 288 289 290 291 292 293 294 295 296 297 298 299 300 301 302 303 304 305 306 307 308 309 310 311 312 313 314 315 316 317 318 319 320 321 322 323 324 325 326 327 328 329 330 331 332 333 334 335 336 337 338 339 340 341 342 343 344 345 346 347 348 349 350 351 352 353 354 355 356 357 358 359 360 361 362 363 364 365 366 367 368 369 370 371 372 373 374 375 376 377 378 379 380 381 382 383 384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408 409 410 411 412 413 414 415 416 417 418 419 420 421 422 423 424 425 426 427 428 429 430 431 432 433 434 435 436 437 438 439 440 441 442 443 444 445 446 447 448 449 450 451 452 453 454 455 456 457 458 459 460 461 462 463 464 465 466 467 468 469 470 471 472 473 474 475 476 477 478 479 480 481 482 483 484 485 486 487 488 489 490 491 492 493 494 495 496 497 498 499 500 501 502 503 504 505 506 507 508 509 510 511 512 513 514 515 516 517 518 519 520 521 522 523 524 525 526 527 528 529 530 531 532 533 534 535 536 537 538 539 540 541 542 543 544 545 546 547 548 549 550 551 552 553 554 555 556 557 558 559 560 561 562 563 564 565 566 567 568 569 570 571 572 573 574 575 576 577 578 579 580 581 582 583 584 585 586 587 588 589 590 591 592 593 594 595 596 597 598 599 600 601 602 603 604 605 606 607 608 609 610 611 612 613 614 615 616 617 618 619 620 621 622 623 624 625 626 627 628 629 630 631 632 633 634 635 636 637 638 639 640 641 642 643 644 645 646 647 648 649 650 651 652 653 654 655 656 657 658 659 660 661 662 663 664 665 666 667 668 669 670 671 672 673 674 675 676 677 678 679 680 681 682 683 684 685 686 687 688 689 690 691 692 693 694 695 696 697 698 699 700 701 702 703 704 705 706 707 708 709 710 711 712 713 714 715 716 717 718 719 720 721 722 723 724 725 726 727 728 729 730 731 732 733 734 735 736 737 738 739 740 741 742 743 744 745 746 747 748 749 750 751 752 753 754 755 756 757 758 759 760 761 762 763 764 765 766 767 768 769 770 771 772 773 774 775 776 777 778 779 780 781 782 783 784 785 786 787 788 789 790 791 792 793 794 795 796 797 798 799 800 801 802 803 804 805 806 807 808 809 810 811 812 813 814 815 816 817 818 819 820 821 822 823 824 825 826 827 828 829 830 831 832 833 834 835 836 837 838 839 840 841 842 843 844 845 846 847 848 849 850 851 852 853 854 855 856 857 858 859 860 861 862 863 864 865 866 867 868 869 870 871 872 873 874 875 876 877 878 879 880 881 882 883 884 885 886 887 888 889 890 891 892 893 894 895 896 897 898 899 900 901 902 903 904 905 906 907 908 909 910 911 912 913 914 915 916 917 918 919 920 921 922 923 924 925 926 927 928 929 930 931 932 933 934 935 936 937 938 939 940 941 942 943 944 945 946 947 948 949 950 951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 968 969 970 971 972 973 974 975 976 977 978 979 980 981 982 983 984 985 986 987 988 989 990 991 992 993 994 995 996 997 998 999 1000 1001 1002 1003 1004 1005 1006 1007 1008 1009 1010 1011 1012 1013 1014 1015 1016 1017 1018 1019 1020 1021 1022 1023 1024 1025 1026 1027 1028 1029 1030 1031 1032 1033 1034 1035 1036 1037 1038 1039 1040 1

[illegible]

2001-2002 150 4 032 (1 611 m) Volume 44 4444

[illegible][illegible]

המחברת מודה כי היא לא יודעת להעריך את חשיבות המידע הזה, ולכן היא לא יכולה להעריך את חשיבות המידע הזה.

[illegible]

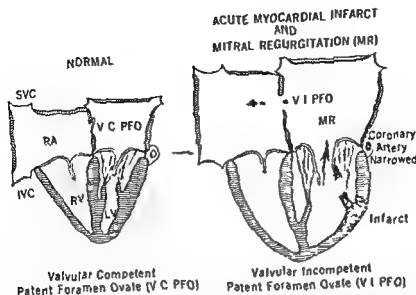


Fig 4 Diagrammatic representation of a valvular-competent patent foramen ovale in contrast to a valvular incompetent patent foramen ovale following myocardial infarction and mitral regurgitation. The left atrial walls have been stretched so that the valve of the patent foramen ovale can no longer cover the defect.

Table I Cardiac catheterization data

| Site | Pressure (mm Hg) | Site | Pressure (mm Hg) |
|------------------|-----------------------------|------------------------|-------------------------------|
| Right atrium | a = 9 v = 21 mean = 9 | Pulmonary artery wedge | a = 25 v = 37 mean = 26 |
| Right ventricle | 54/8 | Left ventricle | 115/29 |
| Pulmonary artery | 62/21 mean = 40 | Brachial artery | 117/92 mean = 108 |

ventricular septal perforation on the basis of a step up in oxygen saturation in the right ventricle unless a left to right shunt at atrial level has been excluded.

Summary

Clinical and necropsy observations are described in a patient who developed a large left to right shunt at atrial level after rupture of a papillary muscle during acute myocardial infarction. Attention is called to the importance of differentiating this combination from rupture of the ventricular septum and from mitral regurgitation

resulting from papillary muscle necrosis with or without rupture.

REFERENCES

1. Tatroles C J, Gault J H, Mason D T and Ross J Jr. Reflux of oxygenated blood into the pulmonary artery in severe mitral regurgitation. *Am Heart J* 15:107 1968.
2. Heikkila J. Mitral incompetence complicating acute myocardial infarction. *Br Heart J* 29:162 1967.
3. Stevenson R R and Turner W J. Rupture of a papillary muscle in the heart as a cause of sudden death. *Bull Johns Hopkins Hosp* 57:235 1935.
4. Davison S. Spontaneous rupture of a papillary muscle in the heart: a report of three cases and



Fig 3 A and B A View of the atrial septum from the right atrium. The fossa ovalis is herniating through the atrial septum. B View of the atrial septum from the left atrium. The fossa ovalis is bulging into the right atrium creating the atrial septal defect. Black silk sutures have closed the surgical atriotomy which is in the interatrial groove.

portion contained about half the chordae tendineae attached to the posteromedial muscle. The atrial septum in the area of the fossa ovalis was stretched and redundant thereby bulging into the right atrium producing a 1.0 cm sized defect which represented a valvular incompetent patent foramen ovale. This defect was closed by suture. The mitral valve was excised and was replaced by a size 19 Hufnagel disc prosthesis. The patient could not be weaned from extracorporeal circulation.

At necropsy (703-376) the left atrium was large and the fossa ovalis was inordinately dilated. A portion of the fossa ovalis herniated into the right atrium enlarging a foramen ovale which had probably been valvular-competent although incompetent before the rupture of the papillary muscle (Fig 3).

Discussion

Papillary muscle rupture as a complication of acute myocardial infarction is rare.² Two large necropsy series discovered only two out of 6,000 and three out of 14,000 cases respectively.² In 1945 Davison³ made the first antemortem diagnosis, and in 1957

Sanders and colleagues¹ compiled 56 cases from the literature and added five new ones. Cederquist and Soderstrom⁴ reported a four year necropsy study in 1964, including 569 patients dying of acute myocardial infarction, from a total of 4,741 cases examined. They found a 0.9 per cent incidence of papillary muscle rupture. Mitral valve replacement has been successful in a number of these patients.¹

The frequency of ventricular septal perforation complicating acute myocardial infarction is probably about 1 per cent.¹¹ Rupture of the ventricular septum classically occurs between 4 and 11 days after acute myocardial infarction and thus overlaps the period during which papillary muscle rupture occurs. The left to right shunt may cause biventricular heart failure.^{10,11}

Mitral valve disease may be associated with left to right shunt at atrial level but usually it is due to rheumatic mitral stenosis and/or insufficiency, rupture of chordae tendineae, or congenital mitral regurgitation.¹⁴ Following acute myocardial infarction and papillary muscle rupture, our patient developed such a shunt from dilation of the left atrium so that the foramen ovale which previously had been valvular competent although patent was stretched and became valvular incompetent (Fig 4). Cases of papillary muscle rupture previously reported have not demonstrated interatrial shunting even though shunts were specifically sought.^{15,16} In rupture of chordae tendineae, however, either spontaneously or after infective endocarditis, interatrial shunting has occurred.^{17,18} Since about 20 to 25 per cent of adults studied at necropsy have valvular-competent but patent foramen ovals,¹⁹ as did this patient it is possible that an interatrial communication might follow acute severe mitral regurgitation of any etiology.

Thus the finding of oxygenated blood in the main pulmonary arteries in patients with acute myocardial infarction and newly acquired systolic precordial murmurs may be due to left to right shunting at atrial level from stretching of the membrane covering the fossa ovalis or from reflux through the capillary bed, or through a shunt at ventricular level. Care should be taken not to diagnose the presence of a

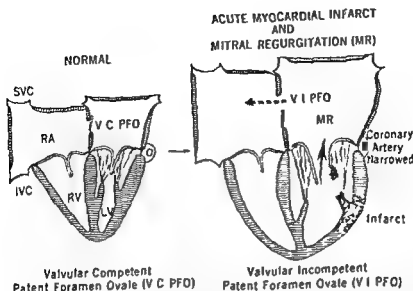


Fig 4 Diagrammatic representation of a valvular competent patent foramen ovale in contrast to a valvular incompetent patent foramen ovale following myocardial infarction and mitral regurgitation. The left atrial wall have been stretched so that the valve of the patent foramen ovale can no longer cover the defect.

Table I Cardiac catheterization data

| Side | Pressure (mm Hg) | Side | Pressure (mm Hg) |
|------------------|-----------------------------|------------------------|-------------------------------|
| Right atrium | s = 9 v = 21 mean = 9 | Pulmonary artery wedge | s = 25 v = 57 mean = 26 |
| Right ventricle | 54/8 | Left ventricle | 115/29 |
| Pulmonary artery | 62/21 mean = 40 | Brachial artery | 117/91 mean = 108 |

ventricular septal perforation on the basis of a step up in oxygen saturation in the right ventricle unless a left to right shunt at atrial level has been excluded.

Summary

Clinical and necropsy observations are described in a patient who developed a large left to right shunt at atrial level after rupture of a papillary muscle during acute myocardial infarction. Attention is called to the importance of differentiating this combination from rupture of the ventricular septum and from mitral regurgitation

resulting from papillary muscle necrosis with or without rupture.

REFERENCES

1. Tatroles C J, Gault J H, Mason D T and Ross J Jr. Reflux of oxygenated blood into the pulmonary artery in severe mitral regurgitation. *Am Heart J* 73: 107, 1968.
2. Heikkila J. Mitral incompetence complicating acute myocardial infarction. *Br Heart J* 29: 167, 1967.
3. Stevenson R R and Turner W J. Rupture of a papillary muscle in the heart as a cause of sudden death. *Bull Johns Hopkins Hosp* 57: 235, 1935.
4. Davison S. Spontaneous rupture of a papillary muscle in the heart: a report of three cases and

- a review of the literature, *J Mt Sinai Hosp N Y* 14:941 1948
- 5 Sanders R J, Neuburger K T and Ravin A Rupture of papillary muscles Occurrence of rupture of posterior muscle in posterior myocardial infarction *Dis Chest* 31:316 1957
- 6 Cederquist L and Soderstrom J Papillary muscle rupture in myocardial infarction a study based upon an autopsy material *Acta Med Scand* 176:287 1964
- 7 Morrow A G, Cohen L S, Roberts W C, Braunwald N S and Braunwald E Severe mitral regurgitation following acute myocardial infarction and ruptured papillary muscle hemodynamic findings and results of operative treatment in four patients *Circulation* 27 and 28 (Suppl 11):124 1963
- 8 Bond V F, Welfare C R, Lide T N and McMillan R I Perforation of the interventricular septum following myocardial infarction *Ann Intern Med* 38:706 1953
- 9 London R L and London S B Rupture of the heart a critical analysis of 47 consecutive autopsy cases *Circulation* 31:707 1965
- 10 Holloway D H, Whalen R F and McIntosh H D Systolic murmur developing after myocardial ischemia or infarction *JAMA* 191:92 1965
- 11 Selzer A, Gerbode F and Kerth W J Clinical hemodynamic and surgical considerations of rupture of the ventricular septum after myocardial infarction *Am Heart J* 78:598 1969
- 12 Demos N J, Gerard F, Sabey A, Ladusky R, Timmes J J and Torruelli J M Coexistence of mitral valve disease and atrial septal defect *J Cardiovasc Surg* 9:278 1968
- 13 Espino-Vela J Rheumatic heart disease associated with atrial septal defect Clinical and pathologic study of twelve cases of Lutenbichers syndrome *Am Heart J* 57:185 1959
- 14 Koss J Jr, Braunwald E, Mason D T, Braunwald N S and Morrow A G Interatrial communication and left atrial hypertension—A cause of continuous murmur *Circulation* 28:853 1963
- 15 Austen W G, Sokol D M, DeSanctis R W and Sanders C A Surgical treatment of papillary muscle rupture complicating myocardial infarction *N Engl J Med* 278:1137 1968
- 16 Breneman G M and Drake E H Ruptured papillary muscle following myocardial infarction with long survival Report of two cases *Circulation* 28:367 1962
- 17 Burchell J B Possibly unrecognized forms of heart disease *Circulation* 28:1153 1963
- 18 Marshall K J and Warden H E Mitral valve disease complicated by left to right shunt at atrial level *Circulation* 29:432 1964
- 19 Edwards J E Pathology of the heart Springfield Ill 1960 Charles C Thomas Publisher p 260

New approach in treatment of cardiac decompensation in U S S R

Ernst Simonson MD

Reuben Berman MD

Minneapolis Minn

The routine treatment of cardiac decompensation with cardiac glycosides and diuretics has not changed essentially during the past decades. It is common experience that the therapeutic effect decreases with the severity of decompensation and the frequent side effects of digitalization are disturbing. In some individual patients the threshold between beneficial and side effects of digitalis therapy is narrow. In the United States and Great Britain considerable work has been done to improve diuretic therapy to lessen side effects and toxic effects and to produce diuresis in intractable failure.

In Russia recent studies in management of heart failure have been aimed at biochemical reactions in the heart muscle itself. Maslyuk and associates¹ and Hlykov² working simultaneously but probably independently have attempted to develop a new approach for the treatment of heart failure based on Meerson's pioneering work^{3,4} on experimental ventricular hypertrophy and decompensation including hemodynamic, macroscopic and microscopic structural changes, biochemical changes and energetics changes of the ECG and changes in the autonomic and central nervous system (cardiovascular and conditioned reflexes, EEG). Meerson's earlier results published in two mono-

graphs^{5,6} and numerous articles were reviewed ten years ago.⁷ Already at that time the material (500 rabbits, 50 dogs with experimental aortic stenosis) was impressive and has been considerably enlarged since then. Biochemical changes were and are the central focus of Meerson's investigations.

During the development of experimental left ventricular hypertrophy synthesis of protein and amino acids is increased and declines in the terminal decompensation. The increase of ATPase during development of hypertrophy indicating improved capacity of myosin for dephosphorylation of ATP was considered by Meerson as one of the principal compensatory factors enabling the myocardium to cope with the increased load. As decrease in the later phases of hypertrophy was found to signal approaching cardiac decompensation Meerson assumed that donors of free SH groups (cysteine, methionine) may counteract the decrease of ATPase in heart failure and thus exert a beneficial effect and this was experimentally confirmed. Administration of a combination of methionine, vitamin B12 and ATP in experimental aortic stenosis in rabbits counteracted the terminal decrease of ATPase and DNA. Meerson's work was paralleled by Western investigators^{8,9} however refer-

From the M U S I H I I I Z I S P k A on M n capol M 55404

II pport d by g a t H C 470103 from the N I al M art a d L g l at t

Reel ed f pub l car Oct 11 1972

R p t rego at t Ernt's Sem so MD Mount S al f pial 2215 P rk A e Mline pol 2, M 55404

ences to Western literature are kept to a minimum in the limited scope of this review. Meerson⁹ has considerably expanded the scope of his work, including various types of ventricular hypertrophy (in arterial hypertension, valvular disease, coronary insufficiency, congenital heart disease and toxic cardiac damage) and found the underlying mechanism quite similar. This was true also for the response to environmental stresses (hypoxia, cold). Meerson⁹ recently summarized his work and proposed a general hypothesis of the genesis of cardiac hypertrophy and terminal decompensation.

Of the numerous results, two are particularly pertinent as a basis for the development of the new therapeutic approaches: (1) inhibition of the increase of synthesis of proteins and nucleic acids prevents the development of compensatory hypertrophy in experimental left ventricular hypertrophy, and (2) administration of inhibitors of protein and nucleic acid synthesis in animals with developed ventricular hypertrophy produces rapid cardiac decompensation.¹⁰

In both studies,^{1,2} the treatment with preparations stimulating protein synthesis supplemented the conventional treatment with cardiac glycosides. Meerson and colleagues¹¹ and Klykov¹² have shown that the cardiotonic effect of cardiac glycosides was increased by a combination with preparations activating the synthesis of nucleic acids: particularly vitamin B12, folic acid, pantothenic calcium, and SH donors, were found to be effective in preventing or ameliorating strophantin intoxication (Klykov,¹² Maslyuk and Fedorova¹³).

The purpose and design of both studies was similar, but there were some differences in sample size, functions tested, evaluation and preparations used.

Preparations used and rationale

Maslyuk and associates¹ investigated the effect of various preparations in three treatment groups supplementing cardiac glycosides and diuretics.

Treatment Group A Potassium orotate, 0.5 Gm, 3 times daily, 1 to 1½ hours before meals. Orotic acid is a precursor in pyrimidine synthesis and is thought to stimulate the synthesis of proteins.

Treatment Group B ATP, 1 ml (10 mg), 2 times daily intramuscular; vitamin B12,

subcutaneous injection 200 mcg daily, methionine orally 0.25 Gm, 3 times daily. While only a negligible part of injected ATP enters the cell, the breakdown products of ATP (AMP and ADP) stimulate oxidative phosphorylation in mitochondria. Vitamin B12 stimulates synthesis of thymine, thymidine, and desoxyribose which are precursors for DNA synthesis. Methionine as donor of SH groups may prevent decrease of ATPase activity of myosin.

Treatment Group C Calcium pantothenate 250 mg, 3 times a day, orally or 1 ml of 20 per cent solution 2 times intramuscular. Pantothenic acid contains acetylcoenzyme A involved in carbohydrate and fat metabolism. It is decreased in cardiac decompensation.¹⁴

Control Group Full therapeutic doses of digitalis preparations (orally) and intravenous strophantin in periods of acute right or left ventricular failure.

Klykov¹² used a combination of vitamin B12, 500 mcg daily, folic acid 0.02 Gm, 3 times daily as stimulant and source for formation of high energy phosphates, ATP sodium, 1 ml (10 mg) 2 times daily intramuscular, calcium pantothenate 0.1 to 0.2 Gm 3 times daily, and Unitiol (sodium 2,3-dimercaptopropane sulfonate) as donor of SH groups 5 ml of 5 per cent solution 1 to 2 times daily intramuscular. KCl 3 Gm per day. All patients control and treatment groups received intravenous strophantin 0.5 ml of 0.05 per cent solution, and diuretics (hypothiazid). Patients with rheumatic heart disease received also aspirin.

Samples

In both studies patients with cardiac decompensation of categories IIA, IIB, III were used. This classification* generally used in the U S S R is like other classifi-

*Classification of cardiac decompensation (G. B. Lang)

Phase I Compensated at rest. Symptoms of decompensation only in physical work: dyspnea, excessive tachycardia, decrease of work capacity (fatigue).

II Decompensation at rest (edema, pulmonary congestion).

A Disturbance of hemodynamics comparatively mild; usually responds well to treatment.

B Pronounced stagnation of circulation in organs, however without significant disturbance of their functions. Therapy less effective.

III Terminal or dystrophic decompensation. Severe disturbance of hemodynamics. Stable changes of metabolism and functions (irreversible histological changes). Work capacity completely reduced.

Table 1 Effect of treatment (in per cent) of patients in each category of cardiac decompensation for four groups*

| Effect of treatment (per cent) | Control group | | | Treatment group A | | | Treatment group B | | | Treatment group C | | |
|--------------------------------|---------------|-----|-----|-------------------|-----|-----|-------------------|-----|-----|-------------------|-----|-----|
| | Phase | | | Phase | | | Phase | | | Phase | | |
| | IIA | IIB | III | IIA | IIB | III | IIA | IIB | III | IIA | IIB | III |
| Excellent | 47† | 16 | 4 | 97 | 68 | 19 | 91 | 66 | 32 | 87 | 67 | 56 |
| Satisfactory | 45 | 51 | 30 | 0 | 32 | 56 | 0 | 25 | 41 | 13 | 33 | 33 |
| Fair | 4 | 24 | 32 | 0 | 0 | 25 | 7 | 7 | 21 | 0 | 0 | 11 |
| Absent | 1 | 9 | 34 | 3 | 0 | 0 | 2 | 2 | 6 | 0 | 0 | 0 |

* group (1) control (calcium + glycoside and diuretics) (2) treatment group A (calcium in addition to glycoside and diuretics) (3) treatment group B (ATP + calcium + thionine in addition to glycoside and diuretics) (4) treatment group C (calcium + thionine + diuretics) (5) treatment group D (calcium + glycoside + diuretics) († Data are tabulated in the information contained in Fig. 1 Maslyuk and colleagues).

cations somewhat arbitrary and probably involves considerable observer variation. However in the hand of the same observers it provides probably a fairly reliable separation as to the degree of decompensation. There is no doubt that these stages reflect progressive phases of decompensation. Maslyuk and colleagues sample¹ of 500 patients was subdivided into four groups: the control group of 251 patients receiving digitalis and diuretics and three treatment groups—A, B and C (together 249 patients)—receiving different preparations as listed above in addition to digitalis and diuretics. The number of patients in the three categories (IIA, IIB, III) are not listed but control and treatment groups were compared for the same category of decompensation.

Klykov's sample (48 controls, 48 treatment) was considerably smaller but control and treatment groups were well matched as to age, sex and phase of decompensation. The average age of the control group was 53 years and that of the treatment group was 52.5 years. There is no information how well the 4 groups in Maslyuk and associates sample¹ were matched but it may be assumed that they were fairly randomized because of the large sample size. The diet in the control and treatment groups was standardized. Both samples included various etiologies: coronary insufficiency, congenital and acquired valvular disease and rheumatic carditis.

A breakdown into these different types of underlying cardiac disease was given in Klykov's study but was incomplete in Maslyuk and colleagues report¹. It appears that both studies were reasonably well controlled but a double blind design would have been superior. At least the control groups should have been given placebos in addition to the conventional digitalis therapy and both patients and hospital personnel should have been unaware who received the placebos and who received the treatment. It is obvious that this might affect the subjective symptoms rather than objective findings but subjective symptoms are essential for the clinical judgment of therapeutic effects.

Methods

The clinical judgment of therapeutic effects was based on subjective symptoms: decrease of cyanosis and hydration (hydrothorax, pulmonary congestion, ascites, liver size, peripheral edema), decrease of respiratory and heart rate, diuresis etc. in addition to investigation of various hemodynamic functions, ECG and blood chemistry.

Results

Clinical course. Table 1 shows the effect of treatment of the four groups in Maslyuk and colleagues study¹ subdivided as to the phase of decompensation (IIA, IIB and III) in terms of excellent, satisfactory

fair and absent Table I shows a highly favorable effect of the supplemental treatment in groups A, B and C in all phases of decompensation (IIA, IIB, III) as compared to the conventional treatment with cardiac glycosides and diuretics. The increase of excellent results in the most severe category of decompensation (III) particularly in Group C is impressive, as well as the absence of a missing therapeutic effect. All patients showed improvement during one month of treatment with pantothenate calcium. Consequently, under the effect of treatment many patients originally in category III shifted to category IIB or IIA and from IIB to IIA. The need for diuretics decreased significantly for category IIA and IIB in all treatment groups (A to C).

The results in Klykov's study were similar. In the control group the effect of conventional treatment (digitalis, diuretics) was excellent in 14 (29 per cent) satisfactory in 24 (50 per cent) absent in 9 (21 per cent). In the treatment group an excellent effect was noted in 36 (75 per cent) eight of 11 patients originally in category III were reclassified into categories IIB or IIA. Since subjective symptoms were involved in the general judgment of therapeutic effect the results are somewhat questionable in the absence of placebo controls. More specific information about the changes of subjective and objective clinical symptoms of cardiac decompensation would have been preferable. However, it appears to be likely that the considerable improvement is not alone psychogenic. In view of this situation objective criteria are more pertinent for the evaluation of therapeutic effects.

Objective criteria In Maslyuk and associates' study¹ the supplemental treatment reduced the incidence of digitalis intoxication (Table II) in all treatment groups A, B, C—particularly in group B (ATP, vitamin B12, methionine) and category III—and was assumed to be due to the effect of free SH groups. ST depression and T inversion improved to a greater extent in all treatment groups than in the control group which may also be due in part to a reduction of digitalis effect.

Table III shows a condensation of Klykov's results of biochemical and hemo-

dynamic changes. There is a greater increase of free SH groups in blood and serum potassium, and a greater decrease of blood pyruvate in the treatment group than in the control group. The changes in the treatment group are highly significant while the changes in the control group did not reach the level of statistical significance.

In both groups there was a highly significant decrease of venous pressure of general and pulmonary circulation time and of the heart rate, but more so in the treatment group. The statistical significance of the difference in the changes in control versus treatment group was not calculated. Respiratory rate decreased significantly in the treatment group, but not in the control group. There was a significant improvement of the mechanics of systole in the treatment group, in contrast in the control group the changes were slight or absent. Particularly impressive is the dramatically faster speed of the increase of intraventricular pressure in the treatment group. Addition of normal values for the items listed in Table III would have been informative.*

For analysis of changes in the mechanical systole Maslyuk and associates¹ divided the sample into several subgroups for decompensation IIA (1) patients with mitral disease (2) patients with aortic and mitral valvular disease for patients in decompensation category IIB and III subgroup (3) patients with mitral disease and subgroup (4) aortic mitral valvular disease. In the control group statistically significant improvement of the parameters of mechanical systole (shortening of isometric phase, lengthening of ejection phase and faster increase of ventricular pressure) were found in subgroups (1) and (2) after treatment with cardiac glycosides. The changes in decompensation categories IIB and III were not statistically significant.

The favorable changes were more pronounced in the treatment groups. In the treatment group A (rotate potassium) significant improvement of systolic parameters were found in subgroups (1) to (3)—

*For normal values of ventricular ejection time and isovolumic contraction time (corresponding approximately to isometric phase) we refer to Aro and Vetter.¹²

Table II *Effect of treatment on incidence of digitalis intoxication**

| Phase | Control group | | | Treatment group A | | | Treatment group B | | | Treatment group C | | |
|---|---------------|-----|-----|-------------------|-----|-----|-------------------|-----|-----|-------------------|-----|-----|
| | IIA | IIB | III | IIA | IIB | III | IIA | IIB | III | IIA | IIB | III |
| Incidence (%) of pronounced bradycardia | 23 | 37 | 45 | 7 | 9 | 31 | 15 | 8 | 12 | 10 | 14 | 21 |
| Frequent PVCs (br emmy) | 13 | 17 | 50 | 0 | 6 | 13 | 3 | 4 | 12 | 10 | 14 | 25 |

Data related to information to be used in Fig. 4. It should be noted that

Table III Changes of hemodynamic functions and blood chemistry in treatment group and control group*

| | Treatment group (N = 43) | | | | | Control group (N = 48) | | | | |
|---|--------------------------|------|-----------------|------|--------|------------------------|------|-----------------|------|--------|
| Blood chemistry and hemodynamics | Before treatment | | After treatment | | | Before treatment | | After treatment | | |
| | M | S.D. | M | C.D. | P | M | S.D. | M | S.D. | P |
| SB groups (m /equ °C) | 56.9 | 1.8 | 65.8 | 1.8 | <0.001 | 57.8 | 2.0 | 67.5 | 1.1 | >0.05 |
| Pyruvic acid (mg %) | 1.66 | 0.11 | 0.94 | 0.04 | <0.001 | 1.94 | 0.19 | 1.41 | 0.13 | >0.05 |
| Potassium (mg %) | 17.5 | 0.6 | 17.6 | 0.9 | <0.001 | 16.5 | 1.1 | 15.1 | 1.1 | >0.1 |
| Hemodynamics | | | | | | | | | | |
| Vascular pressure (mm. H ₂ O) | 19 | 13.9 | 133 | 3 | <0.001 | 150 | 9.9 | 125 | 10.0 | <0.01 |
| Circulation times (sec) | 39.1 | 1 | 4.6 | 0.9 | <0.001 | 32.8 | 1.8 | 71.4 | 1.7 | <0.001 |
| Pulmonary circulation time (sec) | 1.9 | 0.7 | .8" | 0.3 | <0.001 | 11.9 | 0.5 | 10.7 | 0.4 | <0.001 |
| Heart rate | 68.9 | 9 | 77* | 1.5 | <0.001 | 89.2 | *3 | 80.9 | 1.6 | <0.01 |
| Perfusion rate | 77.7 | 0.7 | 90.7 | 0.5 | <0.001 | 70 | 0.5 | 50 | 0.9 | >0.20 |
| Isoelectric contraction phase (msec.) | 51 | 3.0 | ?? | *0 | <0.001 | 49 | 3 | 49 | ? | >0.5 |
| Ejection period (msec) | 17 | 5.0 | 49 | 5.0 | <0.001 | 409 | 6 | 214 | 8 | >0.5 |
| Initial speed of increase of intraventricular press. (mm./Hg/sec) | 1.493 | 31 | 4.167 | 337 | <0.001 | 1.6.3 | 14* | 1.574 | 73 | <0.05 |

*The t test up=1 and AIP L job var= B121 dds to t opha e and dures conz lgr p= str ph i u d
d reusa D t re co d med fom Tabl 1 to 11 Klyk
Abbde all no bi = mean al y SD = m d d d Tasi

ie including decompensation categories IIB and III. In treatment group B (ATP etc.) significant improvement was found in all subgroups (1) to (4). The difference of the changes between control and treatment group II was statistically significant for all subgroups (1) to (4). The changes of electrocardiographic indices also favored the treatment groups.

Table IV shows a summary of the results of the parameters in per cent of inter-

dence of improvement no changes and deterioration. The effect of the supplementary treatment is most obvious in the more advanced categories of decompensation IIB and III—i.e. in conditions with about 45 per cent of failure of the conventional therapy with cardiac glycosides.

In patients with decompensation category IIA hemodynamic indices improved in controls as well as in treatment groups without significant differences between con-

fair and absent Table I shows a highly favorable effect of the supplemental treatment in groups A, B, and C in all phases of decompensation (IIA, IIB, III) as compared to the conventional treatment with cardiac glycosides and diuretics. The increase of excellent results in the most severe category of decompensation (III) particularly in Group C is impressive, as well as the absence of a missing therapeutic effect. All patients showed improvement during one month of treatment with pantothenate calcium. Consequently, under the effect of treatment many patients originally in category III shifted to category IIB or IIA and from IIB to IIA. The need for diuretics decreased significantly for category IIA and IIB in all treatment groups (A to C).

The results in Klykov's study were similar. In the control group the effect of conventional treatment (digitalis diuretics) was excellent in 14 (29 per cent), satisfactory in 24 (50 per cent), absent in 9 (21 per cent). In the treatment group an excellent effect was noted in 36 (75 per cent), eight of 11 patients originally in category III were reclassified into categories IIB or IIA. Since subjective symptoms were involved in the general judgment of therapeutic effect, the results are somewhat questionable in the absence of placebo controls. More specific information about the changes of subjective and objective clinical symptoms of cardiac decompensation would have been preferable. However, it appears to be likely that the considerable improvement is not alone psychogenic. In view of this situation, objective criteria are more pertinent for the evaluation of therapeutic effects.

Objective criteria. In Maslyuk and associates' study,¹ the supplemental treatment reduced the incidence of digitalis intoxication (Table II) in all treatment groups A, B, C—particularly in group B (ATP, vitamin B12, methionine) and category III—and was assumed to be due to the effect of free SH groups. ST depression and T inversion improved to a greater extent in all treatment groups than in the control group, which may also be due in part to a reduction of digitalis effect.

Table III shows a condensation of Klykov's results of biochemical and hemo-

dynamic changes. There is a greater increase of free SH groups in blood and serum potassium, and a greater decrease of blood pyruvate in the treatment group than in the control group. The changes in the treatment group are highly significant, while the changes in the control group did not reach the level of statistical significance.

In both groups there was a highly significant decrease of venous pressure of general and pulmonary circulation time, and of the heart rate, but more so in the treatment group. The statistical significance of the difference in the changes in control versus treatment group was not calculated. Respiratory rate decreased significantly in the treatment group, but not in the control group. There was a significant improvement of the mechanics of systole in the treatment group; in contrast, in the control group the changes were slight or absent. Particularly impressive is the dramatically faster speed of the increase of intraventricular pressure in the treatment group. Addition of normal values for the items listed in Table III would have been informative.*

For analysis of changes in the mechanical systole, Maslyuk and associates¹ divided the sample into several subgroups for decompensation: IIA (1) patients with mitral disease, (2) patients with aortic and mitral valvular disease, for patients in decompensation category IIB and III: subgroup (3) patients with mitral disease and subgroup (4) aortic mitral valvular disease. In the control group, statistically significant improvement of the parameters of mechanical systole (shortening of isometric phase, lengthening of ejection phase, and faster increase of ventricular pressure) were found in subgroups (1) and (2) after treatment with cardiac glycosides. The changes in decompensation categories IIB and III were not statistically significant.

The favorable changes were more pronounced in the treatment groups. In the treatment group A (orotate potassium), significant improvement of systolic parameters were found in subgroups (1) to (3)—

*For normal values of ventricular ejection time and isovolumic contraction time (correspond to approximately 150 ml) refer to Aoyama¹⁰ and Wei¹¹.

- 4 Meerson F Z Problems of pathophysiology of cardiac decompensation (Voprosy patofiziologii nedostatochnost serdtsa Russ) Moscow 1962 Akad Med Nauk p 75
- 5 Simonson E and Lieberman A Russian research on cardiac compensation and decompensation AM HEART J 6:1687 1963
- 6 Gorlin R Recent conceptual advances in congestive heart failure J A M A 179:441 1962
- 7 Bing R J Metabolic activity of the intact heart Am J Med 31:679 1961
- 8 Thauer R Stoffwechsel des Herzmuskels Dtsch G Kreislaufforsch 27:356 1961
- 9 Meerson F Z Development of a present concept of the mechanism of cardiac hypertrophy (Russ) Kardiologia 12(4) 5 1972
- 10 Pshennikova M G Meerson F Z and Teraev N G The influence of folic and orotic acids and actinomycin on the contractile function of the myocardium in hyperfunction of the heart Kardiologia 6(4) 11 1966
- 11 Meerson F Z Pshennikova M G and Pogocyan L A Mechanism of cardiogenic effect of cardiac glycosides (Mekhanizm kardiogeneskogo deistviia serdechnikh glikosidov Russ) Leningrad 1968 Medgiz
- 12 Klykov N V The use of calcium pantothenate in the treatment of chronic cardiac insufficiency Kardiologia 9(2) 130 1969
- 13 Maslyuk V I and Fedorova T A Unitiol in the therapy of intoxication by cardiac glycosides Kardiologia 7(10) 125 1967
- 14 Dedyulin R S Metabolism of pantothenic acid in circulatory insufficiency (Russ) Thesis University of Krasnodar 1967 quoted from (1)
- 15 Weissler A M Harris L C and White G D Left ventricular ejection time index in man J Appl Physiol 18:919 1963
- 16 Aronow W S Bowyer A F and Kaplan M A External isovolumic contraction times and left ventricular ejection time/external isovolumic contraction time ratios at rest and after exercise in coronary heart disease Circulation 43:59 1971

Table IV Changes of parameters of systole during treatment

| Changes in per cent of patients | Decompensation categories | | | | | |
|---------------------------------------|------------------------------------|---|--|------------------------------------|---|--|
| | IIA | | | IIB and III | | |
| | Control (cardiac glycosides) | A Orotate + cardiac glycosides | B ATP B12 etc + cardiac glycosides | Control (cardiac glycosides) | A Orotate + cardiac glycosides | B ATP B12 etc + cardiac glycosides |
| Improvement | 70 10 | 74 3 | 75 0 | 54 6 | 72 7 | 14 2 |
| No change | 21 65 | 25 7 | 25 0 | 11 8 | 22 8 | 20 0 |
| Deterioration | 8 25 | 0 | 0 | 33 6 | 4 5 | 5 8 |

trols and treatment groups. In more severe decompensation (IIB and III), the cardiac minute volume increased 0.4 L. per minute in the control group, 1.0 L. per minute in the treatment group A (difference to controls significant at $p < 0.05$ level), and 1.03 L. per minute ($p < 0.05$) in treatment group B.

Venous pressure decreased in the control group by 39.5 mm H₂O in treatment group A by 61.8 mm (difference to controls $p = 0.01$) and in treatment group B by 80.3 mm ($p < 0.05$). Circulation time decreased in controls 3.4 seconds in treatment group B 8 mm (difference to controls of 4.6 $p < 0.01$) and in treatment group C 6.6 mm ($p < 0.02$).

Thus, in hemodynamic indices as well as in the systolic parameters the favorable effect of the supplementary treatment is more evident in severe cardiac decompensation (IIB, III) than in mild decompensation (IIA).

The significantly greater improvement of various objective indices in the treatment groups as compared to the controls substantiates the general judgment of improved clinical course (Table I) and the improvement of subjective symptoms not specifically listed but included in Table I is in all likelihood secondary to the objective improvement of circulation rather than psychogenic.

Conclusions

It appears that the merit of supplementary treatment of cardiac decompensation

with preparations related to protein synthesis is sufficiently documented to hold promise for further exploration.

Summary

Based on Meerson's fundamental work³ on the genesis of ventricular hypertrophy and decompensation, Russian investigators supplemented the conventional therapy of cardiac decompensation with preparations related to the synthesis of protein and nucleic acids. The merit of the supplementary treatment was shown in the clinical course as well as in various objective criteria (blood chemistry, hemodynamics, systolic parameters, ECG). The reported results are highly significant and the method merits trial in other centers.

We wish to thank Dr. R. M. Gabrielson of Warner Chilcott Laboratories, Morris Plains, N. J. for information about the composition of Unitol and Dr. N. V. Klykov for supplemental information.

REFERENCES

1. Maslyuk V. I., Popov V. G., Popova T. A. and Istvatsev V. P. The treatment of cardiac insufficiency with cardiac glycosides in a complex influencing the synthesis of nucleic acids and energy formation (Russ.) *Kardiologia* 12(1):45, 1972.
2. Klykov N. V. The results of complex treatment of patients with chronic circulatory insufficiency with strophantin, ATP, vitamin B12, folic acid, calcium pantothenate and Unitol (Russ.) *Kardiologia* 12(1):126, 1972.
3. Meerson F. Z. Compensatory cardiac hyperfunction and decompensation (*Kompensatsionnaya i giperfunktsionnaya nedostatochnost serdtsa* (Russ.)) Moscow 1960, Akad. Med. Nauk, p. 258.

Different authors have defined sudden death in coronary heart disease in different ways. In order to make international comparisons concerning sudden non-traumatic death possible, an international expert group agreed in 1970 on the following definition: Sudden unexpected natural death is defined as death occurring immediately or within an estimated period of time of 24 hours after the onset of acute objective or subjective symptoms.¹⁰ Some authors have used 1 hour instead of 24 hours as the criterion.¹¹

Sudden death in acute coronary heart disease

It must be emphasized that not all cases of sudden death, especially when the 24-hour interval is used, are caused by heart disease. Subarachnoid bleeding, other major sudden hemorrhages, massive infections, and pulmonary embolism can also lead to death within such a short interval of time. Most persons who die suddenly, however, do so because of heart conditions. Kuller⁴ reported a study in which 32 per cent of all deaths selected by random sampling were sudden. Sixty per cent of these were considered to be due to acute coronary heart disease.⁴ Spain and co-workers¹⁴ performed a similar investigation in New York and found that a high percentage of sudden deaths was due to heart diseases.

The prerequisite for a scientific analysis of cases of sudden death is a high incidence of autopsy. In Gothenburg, Sweden, the incidence of autopsy is over 90 per cent both in individuals who die in hospital and those who die outside hospital. Forty-five per cent of all patients over 65 years of age who suffered acute myocardial infarction during one year died before arriving in hospital. Under the age of 65 years, the corresponding figure was 25 per cent.¹⁵

Approximately half of all sudden deaths are thus due to coronary heart disease. It must, however, be emphasized that a large part of the patients who die suddenly do not reveal signs of myocardial infarction during autopsy but do exhibit advanced coronary disease. The latter patients often reveal scars due to previous infarcts.^{16,17} It has been pointed out that it may not be

possible to equate patients who die suddenly without pathological anatomical signs of myocardial infarction with those who die similar deaths and who have developed myocardial necrosis.¹⁷ In the extensive World Health Organization study in coronary heart disease and sudden death, it was found that there was a high incidence of patients who lacked thrombosis in the coronary arteries but had pathologically demonstrable myocardial necrosis; they nevertheless died within one hour of the onset of symptoms.¹⁸ Refined methods of investigation, including histochemical staining of autopsy preparations, may produce other results.¹⁹

From the study in Tecumseh, Michigan, 98 deaths due to coronary disease occurred during the six-year follow-up period, 45 of which occurred within one hour of the onset of symptoms.¹²

In the Framingham study,¹³ 156 persons died before reaching the age of 65 years during the 14-year follow-up period, 120 men and 36 women. One hundred twenty of these 156 deaths were due to coronary disease.

In that study, sudden death was defined as death occurring within one hour of the reappearance of acute symptoms. Two thirds of the 120 deaths in coronary disease occurred outside the hospital. Sixty-two of these were sudden deaths as defined above. Younger persons and men were somewhat overrepresented among those who died suddenly, while older persons or women had symptoms for a longer time. Of the 120 deaths in coronary disease, 53 concerned persons in whom this disease had not previously been diagnosed, while 67 of the patients had had diagnosed coronary heart disease before dying. Of these 49, 49 had suffered myocardial infarction. Fifty-one of the 120 had been investigated not more than two years previously and had been classified as completely free from heart disease. It is probable that investigation nearer the time of death—e.g., at an interval of 6 months—would have revealed heart changes. It is concluded that a very high proportion of the deaths which occur suddenly involve individuals who have not previously had serious signs of heart disease.

In a study of labile angina in Edinburgh

Fundamentals of clinical cardiology

Sudden death Identification of high risk groups

*J Anders Vedin M D
Claes Wilhelmsson M D
Dag Elmfeldt M D
Gosta Tibblin M D
Lars Wilhelmsen, M D
Lars Werko M D
Goteborg Sweden*

A substantial number of sudden coronary deaths occur under circumstances which preclude effective treatment. The present paper surveys background knowledge and tries to identify persons with a high risk of future sudden death in order to make the evaluation of primary and secondary preventive efforts effective.

History

In Western Europe and The United States of America cardiovascular diseases are the predominating causes of death.^{1,2} Approximately 50 per cent of men aged between 50 and 64 years who die die of various cardiovascular diseases. In women up to the age of 65 years the share of the total mortality rate occupied by cardiovascular diseases is less than in men but it rises successively with increasing age until it reaches the same level.² Coronary heart disease is the cause of approximately two thirds of the cardiovascular deaths in men and the cause of about one third of cardiovascular deaths in women aged 55 years. With increasing age, the proportion re-

mains relatively stationary in men but increases in women reaching 55 per cent at the age of 75 years.²

In acute episodes of myocardial infarction most patients die outside the hospital. In various studies it has been reported that between 50 and 70 per cent of all deaths in acute myocardial infarction occur before the patient reaches hospital care.^{3,4}

Only in a very small proportion of the deaths has the deceased taken immediate steps to attempt to establish contact with medical assistance. In a study recently performed in Stockholm this was true for less than 10 per cent of the patients.⁵ The cause of this low figure is partly the short interval between the sudden onset of symptoms and death.

It is also partly due to the patients who after the acute onset of symptoms do not appreciate the gravity of the situation or do not wish to trouble a doctor at inconvenient times (e.g. middle of the night). This so called 'patient delay' is difficult to measure and has varied in different studies.^{6,7}

From the Research Unit of Preventive Cardiology, Department of Internal Medicine I (Head: Prof. L. Werko, M.D.), University of Gothenburg, Sahlgren's Hospital, Göteborg, Sweden.

Received for publication Aug. 21, 1972.

Reprint requests to: Dr. J. Anders Vedin, Department of Internal Medicine I, Sahlgren's Hospital, S-413 45 Göteborg, Sweden.

infarction within one hour of the appearance of symptoms but was not correlated with the 3 week mortality rate

In the Tecumseh population the incidence of systolic or diastolic hypertension high relative weight and hyperglycemia was found to be higher in those dying suddenly than for the population as a whole. Multiple risk factors were especially common in those dying suddenly.¹¹

Premonitory symptoms

Prodromal symptoms were demonstrated in 50 to 70 per cent of the patients with myocardial infarction on their arrival in the hospital.^{10, 12} Among the patients who died suddenly prodromal symptoms are difficult to study. Kuller¹³ found symptoms from the upper abdomen and chest in 33 per cent of those who died suddenly and these symptoms either appeared or became altered during the month immediately prior to death. Kuller and associates¹³ found that 24 per cent of those who died suddenly in coronary disease had visited the physician during the week before dying. The cause of these visits however could not be ascertained. Kuller¹³ and also Lindstrom¹⁴ (Table I) found that half of those who died had visited a doctor during the 2 to 4 weeks immediately before dying.^{14, 15} As has previously been pointed out coronary heart disease commences with sudden death in about one fifth of the cases. Analysis of these deaths however shows that very few individuals were normal before dying. In Kuller's¹³ study from Baltimore 92 had diabetes hypertension cerebral hemorrhage or had visited a physician during the month prior to dying or had been admitted to hospital during the year before dying. In the Tecumseh study only one of the 45 who died suddenly had been free from previous illness risk factors and ECG abnormalities in the first cross sectional investigation.¹¹

Arrhythmias in the population

The lethal mechanism in acute heart death has been considered to be malignant ventricular arrhythmias. The extent of arrhythmias in the population or parts of it has been difficult to assess. In studies of life insurance companies in the USA the mortality risk was reported to be greater

Table I Time between last contact with a doctor and death in Goshenburgh (consecutive cases during three months)

| | No autopsy | Fresh MI | Other heart changes |
|-----------------------------|------------|----------|---------------------|
| 0-2 days (n = 7) | 4 | 2 | 1 |
| 3-14 days (n = 11) | 1 | 4 | 6 |
| The last year (n = 12) | 1 | 4 | 7 |
| No known contact (n = 5) | 2 | 1 | 2 |

MI = myocardial infarction.

when premature contractions were present on the ECG. The increase in risk was small but significant.^{16, 17} Prospective population studies confirm the increased risk of death.^{11, 18, 19} In the Tecumseh study 98 individuals died in coronary disease during the six year follow up period. Forty five of these deaths occurred within one hour of the appearance of symptoms. In all but seven of these disturbances of rhythm or conduction had previously been observed on the resting ECG. Ten of these patients (22 per cent) had premature ventricular contractions at rest long before the terminal episode. In all premature contractions were noted in 165 individuals over 30 years of age and among the 3459 individuals who were free from arrhythmias 35 died suddenly. In the individuals with premature ventricular contractions this is equivalent to a sudden death mortality rate of 61 per 1000 compared to 10 per 1000 amongst the individuals who were free from premature contractions. Of the ten deaths among individuals with premature contractions 7 occurred in persons over 60 years of age. The occurrence of premature ventricular contractions was correlated to the occurrence of the coronary disease. The influence of other cardiovascular risk factors was not checked statistically in this study. However no definite connection between accepted cardiovascular risk factors and the tendency to premature ventricular contractions was found. The authors therefore speculated whether the premature ven-

Fulton and others²⁰ have also studied the question of sudden death. Of 380 patients with symptoms suggesting myocardial ischemia, 167 had what was called "unstable angina, or labile angina." Three of these died suddenly and 23 developed acute myocardial infarction within three months. During the same period there were 79 cases of sudden death within the same area, the majority of whom had not, as far as is known, previously experienced pain in the chest. Approximately half of them had consulted a physician for various complaints but these could not be related to myocardial ischemia during the previous months. In this investigation, it was thus found that sudden death was seldom a consequence of labile angina. This investigation thus shows that labile angina or diffuse pain in the chest cannot be used as a prodromal symptom to sudden death.²⁰

Other attempts have been made to calculate the occurrence of sudden death, such as the first manifestation of previously unknown coronary heart disease in the individual—i.e., the first and only manifestation in individuals previously free from infarction or angina pectoris. Despite varying selection of the populations for investigation and varying diagnostic criteria, most of the authors have quoted a figure between 20 and 25 per cent of sudden death from coronary heart disease. This is probably due to the fact that the majority of deaths occur during the hours immediately following the onset and that variations in the period of observation thus cause only minor additions to the number of deaths.²¹⁻²³

In an investigation from Ontario²⁴ it was concluded that, especially during longer periods of time, sudden death cannot be used as an index of coronary heart disease since sudden death in other diseases can vary considerably at different times. This was especially so when the changes in Ontario between 1901 and 1931 were compared with the changes between 1931 and 1951.

The importance of environmental factors in sudden heart death

External temperature. It is well known that patients with angina pectoris suffer exacerbation in cold weather. It has also

been shown that in patients with coronary disease, the ECG changes are accentuated when working in cold weather compared to when working in warmer surroundings.²⁵ Gorbатов²¹ stated in a review of the literature in 1961 that most authors report higher mortality rates when the external temperature is low while others have not been able to confirm this observation.²¹ Westlund² for example, was not able to confirm this in his studies in Oslo. Rose²² demonstrated correlation between cardiovascular mortality rates and low external temperatures, as Swedish authors have also done.²³ In a Swedish study²³ comprising cases of death outside hospital, it was found that there was a correlation between the external temperature and deaths outside the hospital.

The hardness of water. There are a large number of published observations reporting a connection between soft drinking water and a high mortality rate in coronary heart disease.²⁶⁻²⁸ Data from Great Britain suggest that the observed correlation is not associated with concomitant socioeconomic factors and that calcium or other factors are the substances in the water which appear to be of importance.²⁹⁻³² In the above mentioned studies the deaths have not generally been divided up into sudden deaths and other deaths. Data from Ontario show that differences in the hardness of water are related to differences in the occurrence of sudden death outside the hospital and that these differences correspond exactly to the differences in total coronary mortality.³³⁻³⁵

Characteristics for individuals dying suddenly

Prospective studies have demonstrated that sudden death is more common in men than in women and that the occurrence of sudden deaths increases with increasing age.^{2,26,4,46} The risk factors for sudden death are in general agreement with those for coronary heart diseases generally—i.e., hypertension, hypercholesterolemia, obesity, diabetes, cigarette smoking and physical inactivity.^{2,22,43,49} In the Framingham study hypercholesterolemia and ECG signs of left ventricular hypertrophy were associated with sudden death.²² Cigarette smoking was correlated with death in myocardial

infarction within one hour of the appearance of symptoms but was not correlated with the 3 week mortality rate

In the Tecumseh population the incidence of systolic or diastolic hypertension high relative weight and hyperglycemia was found to be higher in those dying suddenly than for the population as a whole. Multiple risk factors were especially common in those dying suddenly.¹¹

Premonitory symptoms

Prodromal symptoms were demonstrated in 50 to 70 per cent of the patients with myocardial infarction on their arrival in the hospital.¹²⁻¹⁴ Among the patients who died suddenly prodromal symptoms are difficult to study. Kinken¹⁴ found symptoms from the upper abdomen and chest in 33 per cent of those who died suddenly and these symptoms either appeared or became altered during the month immediately prior to death. Kuller and associates¹⁵ found that 24 per cent of those who died suddenly in coronary disease had visited the physician during the week before dying. The cause of these visits however could not be ascertained. Kinken¹⁴ and also Lindstrom¹⁶ (Table I) found that half of those who died had visited a doctor during the 2 to 4 weeks immediately before dying.^{14,16} As has previously been pointed out coronary heart disease commences with sudden death in about one fifth of the cases. Analysis of these deaths however shows that very few individuals were normal before dying. In Kuller's¹⁵ study from Baltimore 92 had diabetes hypertension cerebral hemorrhage or had visited a physician during the month prior to dying or had been admitted to hospital during the year before dying. In the Tecumseh study only one of the 45 who died suddenly had been free from previous illness risk factors and ECG abnormalities in the first cross sectional investigation.¹¹

Arrhythmias in the population

The lethal mechanism in acute heart death has been considered to be malignant ventricular arrhythmias. The extent of arrhythmias in the population or parts of it has been difficult to assess. In studies of life insurance companies in the USA the mortality risk was reported to be greater

Table I Time between last contact with a doctor and death in Gothenburg (consecutive cases during three months)

| | No autopsy | Fresh MI | Other heart changes |
|-----------------------------|------------|----------|---------------------|
| 0-2 days (n = 7) | 4 | 2 | 1 |
| 3-14 days (n = 11) | 1 | 4 | 6 |
| The last year (n = 12) | 1 | 4 | 7 |
| No known contact (n = 5) | 2 | 1 | 2 |

MI = myocardial infarction.

when premature contractions were present on the ECG. The increase in risk was small but significant.¹⁷⁻¹⁹ Prospective population studies confirm the increased risk of death.^{11,20-22} In the Tecumseh study 98 individuals died in coronary disease during the six year follow up period. Forty five of these deaths occurred within one hour of the appearance of symptoms. In all but seven of these disturbances of rhythm or conduction had previously been observed on the resting ECG. Ten of these patients (22 per cent) had premature ventricular contractions at rest long before the terminal episode. In all premature contractions were noted in 165 individuals over 30 years of age and among the 3459 individuals who were free from arrhythmias 35 died suddenly. In the individuals with premature ventricular contractions this is equivalent to a sudden death mortality rate of 61 per 1000 compared to 10 per 1000 amongst the individuals who were free from premature contractions. Of the ten deaths among individuals with premature contractions 7 occurred in persons over 60 years of age. The occurrence of premature ventricular contractions was correlated to the occurrence of the coronary disease. The influence of other cardiovascular risk factors was not checked statistically in this study. However no definite connection between accepted cardiovascular risk factors and the tendency to premature ventricular contractions was found. The authors therefore speculated whether the premature ven-

tricular contractions themselves imply an increased risk of sudden death or only constitute another sign of other predisposing factors.¹¹ Pell and d'Alonzo¹² found in another study among the employees of a large company that the very early mortality rate in coronary disease was associated with older age, previous hypertension, and ECG anomalies such as disturbances of conduction and premature ventricular contractions.

In the 'seven countries' study by Blackburn, Taylor, and Keyes¹³ it was found that the occurrence of premature ventricular contractions at rest detected in the general population (> 10 per cent of the recorded beats) was correlated to the incidence of coronary disease and the mortality rates recorded during the follow up period. In this study six concomitant factors were investigated (age, skinfold, systolic blood pressure, serum cholesterol, smoking, and the level of physical activity). Premature contractions after working (3 min step test) were not of the same prognostic significance as premature contractions recorded at rest, but the analysis was not complete. Hinkle and colleagues¹⁴ used portable ECG recorders in a study of employees of a company. During a recording period of six hours the disturbances of rhythm and conduction were recorded in not less than 92.6 per cent of the cases. The most common arrhythmia consisted of premature ventricular contractions which were found in 62.2 per cent. In this study too, the occurrence of ventricular arrhythmias was correlated to findings of coronary heart disease. The study confirmed without standardization for concomitant risk factors that the finding of premature ventricular contractions was associated with future coronary death. The connection became more pronounced with increasing occurrence of ventricular contractions. In Gothenburg too, we have confirmed that there is a connection between premature ventricular contractions and signs of coronary heart disease.¹⁵

None of the studies so far can be considered to have solved the question whether premature contractions in the population are directly or indirectly associated with the phenomenon of sudden death.

Studies on infarction patients

Since approximately half of all sudden deaths are directly related to coronary disease, and since more than half of all coronary patients will die suddenly without knowledge of their danger, it is evident that the most effective discriminatory factor concerning sudden death is previous myocardial infarction.

Premonitory symptoms in patients with previous myocardial infarction

Progressive angina pectoris, so-called "unstable angina," has attracted attention as a premonitory symptom.¹⁶ It is possible that this symptom is more common prior to infarction or death, but angina pectoris itself is a variable symptom. Many symptoms noted during the preinfarction period are also common in the general population and the specific discriminatory value of the symptom is not known and at present cannot be calculated.

In Gothenburg we have at present a group of more than 500 infarction patients below 67 years of age and constituting all hospitalized infarction patients in Gothenburg between 1968 and 1970.

Premonitory symptoms in patients with previous myocardial infarction have so far not been specifically studied in the post myocardial infarction clinic in Gothenburg. However, patients have been in very close contact with the clinic and have received specific instructions to consult the physicians or nurses if any symptoms causing them the least concern should occur. Almost all of the 38 patients who died suddenly had been in contact with the clinic within one month before dying. On retrospective investigation no obvious prodromal signs were detected.¹⁴

At the moment most of the prodromal symptoms are vague, not only for the patients but also for the medical profession and therefore fail to activate adequate measures. There is no doubt that at the moment extremely little is known about the treatment of unstable angina.

High risk groups among patients with previous infarction. A prognostic index has been established in order to define groups of infarction patients with varying prognosis.¹⁵ Simple factors recorded during

Sub-group I No extensive cardiac damage

Sub-group II Mechanical damage to myocardium

- 1) Rel heart volume $\left\{ \begin{array}{l} \sigma > 450 \text{ ml/m}^2 \text{ BSA} \\ \rho > 400 \text{ ml/m}^2 \text{ BSA} \end{array} \right.$
- 2) GPT > 40 U during first 3 days
- 3) BT > 38°C
- 4) Transient atrial flutter and fibrillation

Sub-group III Electrical cardiac damage

- 1) VPB frequency > 5/min
- 2) Ventricular tachycardia or ventricular fibrillation
- 3) AV-blocks $\left\{ \begin{array}{l} a \text{ PQ} > 0.24 \text{ s} \\ b \text{ Type II} \\ c \text{ Type III} \end{array} \right.$

Sub-group IV Combined electro-mechanical damage (Sub-groups II or III)

Fig 1 Criteria for homogeneous sub group allocation of patients 57 to 67 years of age with myocardial infarction. Explanation of the schematic as follows: Sub group I: no extensive cardiac damage. Sub group II: mechanical damage to myocardium (1) relative heart volume (men = > 450 ml per square meter of body surface area, women = > 400 ml per square meter of body surface area), (2) glutamic pyruvic transaminase > 40 units during first three days, (3) body temperature > 38°C during first three days, and (4) transient atrial flutter and fibrillation. Sub group III: electrical cardiac damage (1) ventricular premature beat frequency > 5 per minute, (2) ventricular tachycardia or ventricular fibrillation, and (3) AV blocks (a) PQ interval > 0.24 sec, (b) Type II, and (c) Type III. Sub group IV: combined electro-mechanical damage (sub groups II or III).

every infarction patient's stay in hospital enable a prognosis to be established (Fig 1).^{48, 49} The criteria for allocation of patients to subgroups with regard to risk have been obtained from a previous study on patients in a post myocardial infarction clinic in Gothenburg.⁴⁸ Each single factor was correlated to death as reinfarction during the first post hospital year. For convenience the criteria were grouped as follows: (1) lack of extensive cardiac damage, (2) mechanical cardiac damage, (3) electrical cardiac damage, and (4) combined electro-mechanical damage to the heart. The presence of isolated criteria is sufficient for allocation to a certain group. The proportional distribution of the infarction patients and the incidence of death and reinfarction in the

various groups is shown in Fig 2. Fig 2 also shows the distribution of mortality rate in the groups after an average follow up time of 7.5 months.⁴⁹

In the present series of patients discharged from the hospital, ventricular arrhythmias occurring during the acute stage are practically always associated with later detected signs of major mechanical myocardial injury (Fig 2). Almost all patients in this series who have had ventricular arrhythmias while in the hospital show signs of myocardial injury.

Arrhythmias in patients with previous infarction. In total populations in which the prevalence of coronary heart diseases has been low, a connection has nevertheless been demonstrated between premature

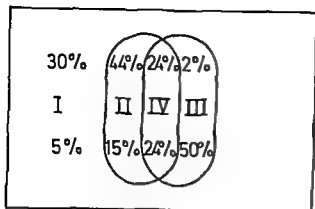


Fig 2 The percentage of patients with myocardial infarction in secondary risk groups (upper register) and the rate of reinfarction and mortality (bottom register). Average follow up period = 7.5 months

ventricular contractions and coronary disease and also in association between previously recorded premature ventricular contractions and sudden death. Furthermore in the group of individuals who have died sudden deaths signs of previous or existing coronary disease have been observed. Among patients discharged from the hospital in Gothenburg after myocardial infarction above 57 years of age 77 per cent of the deaths were sudden and below 57 years of age 46 per cent of the deaths were sudden.⁶⁴

It is well known that acute ventricular arrhythmias are the cause of the high initial mortality rate in ischemic heart disease. Ventricular arrhythmias during the acute stage are correlated to an increase in late mortality rate.⁶⁵ The incidence of premature ventricular contractions after myocardial infarction was associated with an increased risk of mortality as was shown for the first time among the infarction patients of the Coronary Drug Project.⁶⁶

If the period of ECG recording is prolonged 80 per cent of patients have premature ventricular contractions after infarction.⁷⁰⁻⁷¹ This is therefore not an adequately effective discriminatory factor for sudden death after myocardial infarction. The frequency of premature contractions and the circumstances under which arrhythmias occur—e.g., physical work—may be of importance.⁶⁷⁻⁷² Various methods of provoking and grading arrhythmias in the infarction patients may enable the allocation of patients to various risk groups concerning sudden death.

Conclusions

If primary preventive measures taken before the onset of coronary disease are rewarded with success, it may be expected that this will also reduce the occurrence of sudden death.

However it is known that a large number of individuals who later die suddenly have been in contact with a physician during the period immediately prior to dying.

The occurrence of sudden death may be expected to be high in a group of myocardial infarction patients discharged from the hospital. Every effort to prevent or delay sudden death should be directed at this group in the first place.

Our knowledge concerning possible prodromal symptoms and their treatment is at present much too scanty to enable practical recommendations.

In view of the present uncertainty concerning the significance of premature ventricular contractions in the chronic stage it is important that controlled studies be performed concerning the ability of various drugs not only to suppress premature ventricular contractions but also to reduce the occurrence of sudden death.

REFERENCES

1. Stamler J. Cardiovascular diseases in the United States. *Am J Cardiol* 10:319 1967.
2. Vedin J, A. Wilhelmsson C, E. Bolander A, M. and Werko L. Mortality trends in Sweden 1951-1968 with special reference to cardiovascular causes of death. *Acta Med Scand* 515(Suppl 1):76 1971.
3. Kuller L. Sudden and unexpected nontraumatic deaths in adults. *J Chronic Dis* 19:1165 1966.
4. Kuller L, Littenfeld A, and Fisher R. Epidemiological study of sudden and unexpected deaths due to arteriosclerotic heart disease. *Circulation* 34:1056 1966.
5. Kuller L, Littenfeld A, and Fisher R. An epidemiological study of sudden and expected deaths in adults. *Medicine* 46:341 1967.
6. Elmfeldt D, and Wilhelmsson L. Hjärtans färdsmörbidenhet, mortalitet och livslängd. *Läkartidningen* 68:3705 1971.
7. Wiklund B. Medically unattended fatal cases of ischaemic heart disease in a defined population. *Acta Med Scand* 524(Suppl 1):78 1971.
8. Fulton M, Julian D, G. and Oliver M. F. Sudden death and myocardial infarction. *Circulation* 39 and 40(Suppl 4):182 1969.
9. Lindholm B. The ischaemic heart disease register. *Fehr Dubb J* 4(Suppl 111):34 1967.
10. Paul O, and Schatz M. On sudden death (Editorial). *Circulation* 43:7 1971.

- 11 Chiang B N Perlman L V Ostrander L D Jr and Epstein F H Relationship of premature systoles to coronary heart disease and sudden death in the Tecumseh epidemiologic study *Ann Intern Med* 10:1159 1969
- 12 Chiang B N Perlman L V Fulton M Ostrander L D Jr and Epstein F H Predisposing factors in sudden cardiac deaths in Tecumseh Michigan *Circulation* 41 31 1970
- 13 Gordon T and Kannel W B Premature mortality from coronary heart disease The Framingham Study *JAMA* 215:1617 1971
- 14 Spain D M Bradess V A and Mohr C Coronary atherosclerosis as a cause of unexpected and unexplained death An autopsy study from 1949-1959 *JAMA* 174:384 1960
- 15 Fodor J and Tibblin G One year's experience of the ischemic heart disease registry in Gothenburg in ischemic heart disease registers WHO Working Document Geneva 1970 EURO 5010(4) pp 10-14
- 16 Lehman L Lindström B Tibblin G and Elmfeldt D Unattended deaths in heart disease in Göteborg Fifth Annual Meeting in Scheveningen The Netherlands 1971 *Eur Soc Clin Invest* pp 32-33
- 17 Tibblin G The men born in 1913 in Tibblin G Keys A and Werkö L editors Preventive cardiology Stockholm 1971 Almqvist & Wiksell J Wiley & Sons New York
- 18 Hagan A Livaic A M Sternby H and Vihert A M Coronary artery thrombosis and the acute attack of coronary heart disease *Lancet* 11:1199 1968
- 19 Andersen J A Fischer R Hansen B Kaaber K and Thrane L Udvidet hjerteaftopsi Ugeskr Laeger 134 990 1972
- 20 Fulton M Duncan B Lutz W et al Natural history of unstable angina *Lancet* 1 860 1972
- 21 Morris J N Heady M A and Barley R G Coronary heart disease in medical practitioners *Br Med J* 1:503 1952
- 22 Zukel W L Lewis R H Enterline P E Painter R C Ralston L S Fawcett R M Meredith A P and Peterson M A short term community study of the epidemiology of coronary heart disease *Am J Public Health* 49:1630 1959
- 23 Doyle J T Heslin A S Hilleboe H E and Formel P F Early diagnosis of ischemic heart disease *N Engl J Med* 261:1096 1959
- 24 Eisenberg H Feltner W R Payne G H and Haddad C A The epidemiology of coronary heart disease in Middlesex County Connecticut A preliminary report on methodology and the incidence of primary myocardial infarction *J Chronic Dis* 14:221 1961
- 25 Kannel W B Hagan A Dawber T R and Revotskie N Epidemiology of coronary heart disease Implications for the practicing physician *Geriatrics* 17 675 1962
- 26 Pell S and d'Alonzo C A Immediate mortality and five-year survival of employed men with a first myocardial infarction *N Engl J Med* 270 915 1964
- 27 Shapiro S Weinblatt E Frank C W and Sager R V The HIP study of incidence and prognosis of coronary heart disease Preliminary findings on incidence of myocardial infarction and angina *J Chronic Dis* 18:527 1965
- 28 Mathewson F A L Breckton C C Keltie W A and Paul G J The University of Manitoba follow up study A prospective investigation of cardiovascular disease *Can Med Assoc J* 92:947 1965
- 29 Anderson T W and Le Riche W H Ischaemic heart disease and sudden death 1901-61 *Br J Prev Soc Med* 24:1 1970
- 30 Blomqvist G The frank lead exercise electrocardiogram A quantitative study based on averaging technique and digital computer analysis *Acta Med Scand* 178(Suppl 440) 1 1965
- 31 Gorbatow L Die Herzinfarktfräquenz in einem Material der Bevölkerung von Helsinki während der Jahre 1945-1952 *Acta Med Scand* (Suppl 364) 1 1961
- 32 Westlung K Further observations on the incidence of myocardial infarction in Oslo *J Oslo City Hosp* 13:201 1963
- 33 Rose G Cold weather and ischemic heart disease *Br J Prev Soc Med* 20 97 1966
- 34 Hall P Mosezon E Belander H and Selander K Utombustemperatur och mortalitet i hjärt och kärlsjukdomar *Läkartidningen* 67:2141 1970
- 35 Falconer B De primära hjärt aortadödsfallen ettärsmaterial från rattfäklarstation *Läkartidningen* 69:764 1972
- 36 Schroeder H A Relation between mortality from cardiovascular disease and treated water supplies variations in states and 163 largest municipalities of United States *JAMA* 172 1902 1960
- 37 Morris J N Crawford M D and Heady J A Hardness of local water supplies and mortality from cardiovascular disease in county boroughs of England and Wales *Lancet* 1:860 1961
- 38 Björck G Bostrom H and Widstrom A On relationship between water hardness and death rate in cardiovascular diseases *Acta Med Scand* 18 739 1965
- 39 Crawford M D Gardner M J and Morris J N Mortality and hardness of local water supplies *Lancet* 1 827 1968
- 40 Robertson J S Mortality and hardness of water *Lancet* 11:348 1968
- 41 Neri L C Mandel J and Hewitt M Relation between mortality and water hardness in Canada *Lancet* 1:931 1972
- 42 Roberts C J and Lloyd S Association between mortality from ischemic heart disease and rain fall in South Wales and in the County boroughs of England and Wales *Lancet* 1:1091 1972
- 43 Anderson T W Le Riche W H and Machay J S Sudden death and ischemic heart disease correlation with hardness of local water supply *N Engl J Med* 280 805 1969
- 44 Anderson T W and Le Riche W H Sudden death from ischemic heart disease in Ontario

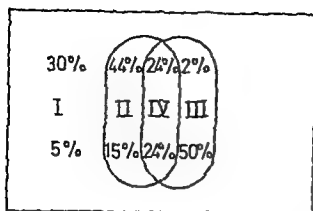


Fig 2 The percentage of patients with myocardial infarction in secondary risk groups (upper register) and the rate of reinfarction and mortality (bottom register). Average follow up period = 7.5 months

ventricular contractions and coronary disease and also an association between previously recorded premature ventricular contractions and sudden death. Furthermore in the group of individuals who have died sudden deaths signs of previous or existing coronary disease have been observed. Among patients discharged from the hospital in Gothenburg after myocardial infarction above 57 years of age 77 per cent of the deaths were sudden and below 57 years of age 46 per cent of the deaths were sudden.¹⁴

It is well known that acute ventricular arrhythmias are the cause of the high initial mortality rate in ischemic heart disease. Ventricular arrhythmias during the acute stage are correlated to an increase in late mortality rate.¹⁵ The incidence of premature ventricular contractions after myocardial infarction was associated with an increased risk of mortality as was shown for the first time among the infarction patients of the Coronary Drug Project.¹⁶

If the period of ECG recording is prolonged, 80 per cent of patients have premature ventricular contractions after infarction.^{10, 11} This is therefore not an adequately effective discriminatory factor for sudden death after myocardial infarction. The frequency of premature contractions and the circumstances under which arrhythmias occur—e.g., physical work—may be of importance.^{12, 13} Various methods of provoking and grading arrhythmias in the infarction patients may enable the allocation of patients to various risk groups concerning sudden death.

Conclusions

If primary preventive measures taken before the onset of coronary disease are rewarded with success, it may be expected that this will also reduce the occurrence of sudden death.

However, it is known that a large number of individuals who later die suddenly have been in contact with a physician during the period immediately prior to dying.

The occurrence of sudden death may be expected to be high in a group of myocardial infarction patients discharged from the hospital. Every effort to prevent or delay sudden death should be directed at this group in the first place.

Our knowledge concerning possible prodromal symptoms and their treatment is at present much too scanty to enable practical recommendations.

In view of the present uncertainty concerning the significance of premature ventricular contractions in the chronic stage it is important that controlled studies be performed concerning the ability of various drugs not only to suppress premature ventricular contractions but also to reduce the occurrence of sudden death.

REFERENCES

1. Stamler J. Cardiovascular diseases in the United States. *Am J Cardiol* 10:319, 1967.
2. Vedin J, A. Wilhelmsson C, E. Bolander A, M. and Werko L. Mortality trends in Sweden 1951-1968 with special reference to cardiovascular causes of death. *Acta Med Scand* 515(Suppl 1):76, 1971.
3. Kuller L. Sudden and unexpected nontraumatic deaths in adults. *J Chronic Dis* 19:1165, 1966.
4. Kuller L, Liliensfeldt A and Fisher R. Epidemiological study of sudden and unexpected deaths due to arteriosclerotic heart disease. *Circulation* 34:1056, 1966.
5. Kuller L, Liliensfeldt A and Fisher R. An epidemiological study of sudden and expected deaths in adults. *Medicine* 46:341, 1967.
6. Elmfeldt D and Wilhelmsson L. Hjärtinfarktombudet mortalitet och invaliditet. *Läkartidningen* 68:3105, 1971.
7. Wiklund B. Medically unattended fatal cases of ischaemic heart disease in a defined population. *Acta Med Scand* 524(Suppl 1):78, 1971.
8. Fulton M, Julian D G and Oliver M F. Sudden death and myocardial infarction. *Circulation* 39 and 40(Suppl 4):182, 1969.
9. Lindholm B. The ischaemic heart disease register. *Pehr Dabb J* 4(Suppl 111):34, 1969.
10. Paul O and Schatz M. On sudden death (Editorial). *Circulation* 43:17, 1971.

Appraisal and reappraisal of cardiac therapy

Edited by Arthur C DeGraff and Julian Frieden

Drug therapy of heart disease in pediatric patients

I Congestive heart failure in infancy and concepts of developmental pharmacology

Sanford N Cohen MD*

Eugenie F Doyle MD**

Monika M Rutkowski MD***

New York NY

Etiologic and hemodynamic features of heart failure in infancy

Congestive heart failure in infancy is a common serious pediatric problem. The younger an infant is when signs and symptoms of cardiac decompensation occur, the worse the prognosis. A variety of disease entities may cause heart failure under one year of age (Table I) but by far the leading causes are congenital cardiovascular malformations. These are present in approximately 8 of every 1,000 live born infants. Thirty per cent of these infants die during the first year of life, most of them in congestive heart failure. Since these malformations may lead to rapid deterioration of the patient's condition, strenuous measures must be employed to treat the cardiac failure, diagnose the cardiac defect accurately, and refer the patient for prompt surgery if the defect is amenable to palliation or correction.

Whatever the etiology of cardiac failure might be, the fundamental hemodynamic

problem is the same: the cardiac output is inadequate for the needs of the body and excessive quantities of blood accumulate in the pulmonary and systemic venous systems. A rise in ventricular end diastolic and atrial filling (mean) pressure is the initial response to a drop in cardiac output. Subsequently, tachycardia, widening of the arteriovenous oxygen difference, and cardiac dilatation occur. The adrenergic nervous system plays an important role in supporting myocardial contractility; depletion of catecholamines is associated with increasing manifestation of congestive heart failure.

The various congenital cardiovascular defects may cause failure by different mechanisms.

Left to right shunts (Group I in Table I). There is a volume overload on one or both ventricles in the presence of a large left to right shunt. In such a situation, ventricular dilatation and tachycardia occur first, followed by gradually progressive ventricular hypertrophy. There is also an increased

* From the Department of Pediatrics, Philadelphia Children's Hospital, Philadelphia, PA 19104.

Received for publication February 12, 1973.

Reprint requests to Dr. Sanford N. Cohen, Associate Professor of Pediatrics, New York University School of Medicine, 550 1st Avenue, New York, NY 10016.

Associate Professor of Pediatrics, Philadelphia Children's Hospital, Philadelphia, PA 19104.

Associate Professor of Pediatrics, Philadelphia Children's Hospital, Philadelphia, PA 19104.

- and its correlation with water hardness and other factors *Can Med Assoc J* 105:155 1971
- 45 Kannel W B, Barry P and Dawber T R Immediate mortality in coronary heart disease The Framingham Study Mexico 1963 *Memorias IV Congress Mundial de Cardiologia* IV B 176
 - 46 Wilhelmsen L, Elmfeldt D, Vedin A, Wilhelmsson C and Tibblin G (In preparation)
 - 47 Morris J N, Herdy J A, Raffle P A II et al Coronary heart disease and physical activity of work *Lancet* II:1111 1963
 - 48 Wilhelmsen L and Tibblin G Physical inactivity and risk of myocardial infarction—The men born in 1913 study in Larsen O A and Malmberg R O editors *Physical fitness and coronary heart disease* Copenhagen 1971 Munksgaard
 - 49 Doyle J T Cigarette smoking: the associated cardiovascular risk *Minn Med* 52:1311 1969
 - 50 Feil H Preliminary print in coronary thrombosis *Am J Med Sci* 193:42 1937
 - 51 Sampson J J and Elaser M The diagnosis of impending acute coronary artery occlusion *Am Heart J* 13:675 1937
 - 52 Solomon H A, Edwards A L and Killip T Prodromata in acute myocardial infarction *Circulation* 40:463 1969
 - 53 Stowers M and Short D Warning symptoms before major myocardial infarction *Br Heart J* 32:833 1970
 - 54 Kivlen L M D thesis University of Oxford 1969
 - 55 Lindstrom B Death from AHD outside hospital *Pehr Dubb J* 4(Suppl III):38 1969
 - 56 Brandon K F, Neill M H and Streeter G C The use of the electrocardiogram in twenty five years of insurance selection *Trans Assoc Life Ins Med Dir Am* 34:143 1950
 - 57 Kirkland H H, Kuessing C E and Lyle A M The evaluation of certain fundamental electrocardiographic patterns in the selection of insurance risks *Trans Assoc Life Ins Med Dir Am* 35:86 1951
 - 58 Lyle A M Coronary disease as an underwriting problem *Trans Soc Actuaries* 15:324 1963
 - 59 Rodstein M, Wollock L and Gubner R S Mortality study of the significance of extra systoles in an insured population *Circulation* 44:617 1971
 - 60 Blackburn H, Taylor H L and Keys A The electrocardiogram in prediction of five-year coronary heart disease incidence among men aged forty through fifty nine *Circulation* 41(Suppl 1):154 1970
 - 61 Hinkle L E Jr, Carver S T and Stevens M The frequency of asymptomatic disturbances of cardiac rhythm and conduction in middle-aged men *Am J Cardiol* 21:65 1969
 - 62 Vedin J A, Wilhelmsson C E, Wilhelmsen L and Bjure J The relationship of rest and exercise induced ectopic beats to other ischemic manifestations and to coronary risk factors *Am J Cardiol* 30:25 1972
 - 63 Fulton M and Julian D G The Edinburgh chest pain study Geneva 1971 WHO working Document Euro 8204 (3) pp 21-22
 - 64 Wilhelmsen L, Elmfeldt D, Tibblin G, Vedin A and Wilhelmsson C Pre-coronary care Geneva 1972 WHO working document Euro 8204 (4)
 - 65 Norris R M, Brandt P W T and Caughey D E A new coronary prognostic index *Lancet* I:274 1969
 - 66 Elmfeldt D and Wilhelmsen L A study of representative post myocardial infarction patients aged 27-55 in Tibblin G, Keys A and Werkö L editors *Preventive cardiology* Stockholm 1972 Almqvist & Wiksell J Wiley & Sons New York
 - 67 Vedin J A and Wilhelmsson C E Evaluation of a post myocardial infarction outpatient clinic in Tibblin G, Keys A and Werkö L editors *Preventive cardiology* Stockholm 1972 Almqvist & Wiksell J Wiley & Sons New York
 - 68 Denborough M A, Lovell R R H, Nester P J and Goble A J Arrhythmias and late sudden death after myocardial infarction *Lancet* I:386 1968
 - 69 The coronary drug project research group (in press 1972)
 - 70 Lown B Arrhythmias and sudden death: Geneva 1971 WHO working document Euro 8204 (3) pp 17-19
 - 71 Lown B and Wolf M Approaches to sudden death from coronary heart disease *Circulation* 44:130 1971
 - 72 Kosonowsky B D, Lown B, Whiting R and Guiney T Occurrence of ventricular arrhythmias with exercise as compared to monitoring *Circulation* 44:826 1971

Appraisal and reappraisal of cardiac therapy

Edited by Arthur C. DeGraff and Julian Frieden

Drug therapy of heart disease in pediatric patients

I Congestive heart failure in infancy and concepts of developmental pharmacology

Sanford N. Cohen, M.D.*

Eugenie F. Doyle, M.D.**

Monika M. Rutkowski, M.D.***

New York, N.Y.

Etiologic and hemodynamic features of heart failure in infancy

Congestive heart failure in infancy is a common serious pediatric problem. The younger an infant is when signs and symptoms of cardiac decompensation occur, the worse the prognosis. A variety of disease entities may cause heart failure under one year of age (Table I) but by far the leading causes are congenital cardiovascular malformations. These are present in approximately 8 of every 1,000 live born infants. Thirty per cent of these infants die during the first year of life, most of them in congestive heart failure. Since these malformations may lead to rapid deterioration of the patient's condition, strenuous measures must be employed to treat the cardiac failure, diagnose the cardiac defect accurately, and refer the patient for prompt surgery if the defect is amenable to palliation or correction.

Whatever the etiology of cardiac failure, might be the fundamental hemodynamic

problem is the same: the cardiac output is inadequate for the needs of the body, and excessive quantities of blood accumulate in the pulmonary and systemic venous systems. A rise in ventricular end-diastolic and atrial filling (mean) pressure is the initial response to a drop in cardiac output. Subsequently, tachycardia, widening of the arteriovenous oxygen difference, and cardiac dilatation occur. The adrenergic nervous system plays an important role in supporting myocardial contractility; depletion of catecholamines is associated with increasing manifestation of congestive heart failure.

The various congenital cardiovascular defects may cause failure by different mechanisms.

Left to right shunts (Group I-A, Table I). There is a volume overload on one or both ventricles in the presence of a large left to right shunt. In such a situation, ventricular dilatation and tachycardia occur first, followed by gradually progressive ventricular hypertrophy. There is also an increased

*From the Department of Pediatrics and Pharmacology, New York University School of Medicine.

Revised 4/10/73, published 4/12/73.

Reprint requests: Dr. Sanford N. Cohen, Associate Professor of Pediatrics, New York University School of Medicine, 550 1st Ave., New York, N.Y. 10016.

Associate Professor of Pediatrics and Pharmacology, New York University School of Medicine, Physical Chemistry Department, Bellevue Hospital, John and Mary R. Mable School of Academic Medicine.

**Professor of Pediatrics and Director of Pediatric Cardiology, New York University Medical Center.

***Fellow in Pediatrics, New York University School of Medicine.

- and its correlation with water hardness and other factors *Can Med Assoc J* 105:155 1971
- 45 Kannel W B Barry P and Dawber T R Immediate mortality in coronary heart disease The Framingham Study Mexico 1963 *Memorias IV Congress Mundial de Cardiologia* IV B 176
 - 46 Wilhelmsson L Elmfeldt D Vedin A Wilhelmsson C and Tibblin G (In preparation)
 - 47 Morris J N Hardy J A Raffle P A H et al Coronary heart disease and physical activity of work *Lancet* III:1111 1953
 - 48 Wilhelmsson L and Tibblin G Physical inactivity and risk of myocardial infarction—The men born in 1913 study in Larsen O A and Malmberg R O editors *Physical fitness and coronary heart disease* Copenhagen 1971 Munksgaard
 - 49 Doyle J T Cigarette smoking the associated cardiovascular risk *Minn Med* 52:1311 1969
 - 50 Feil H Preliminary pain in coronary thrombosis *Am J Med Sci* 193 42 1937
 - 51 Sampson J J and Elmer M The diagnosis of impending acute coronary artery occlusion *Am Heart J* 13:675 1937
 - 52 Solomon H A Edwards A L and Killip T Prodromata in acute myocardial infarction *Circulation* 40 463 1969
 - 53 Stowers M and Short D Warning symptoms before major myocardial infarction *Br Heart J* 32:833 1970
 - 54 Kinnin L M D thesis University of Oxford 1969
 - 55 Lindström B Death from ASHD outside hospital *Pebr Dubb J* 4(Suppl III) 38 1969
 - 56 Brandon K F Neill M H and Streeter G C The use of the electrocardiogram in twenty five years of insurance selection *Trans Assoc Life Ins Med Dir Am* 34:143 1950
 - 57 Kirkland H B Kiessling C E and Lyle A M The evaluation of certain fundamental electrocardiographic patterns in the selection of insurance risks *Trans Assoc Life Ins Med Dir Am* 35 86 1951
 - 58 Lyle A M Coronary disease as an underwriting problem *Trans Soc Actuaries* 15 324 1963
 - 59 Rodstein M Wollock L and Gubner R S Mortality study of the significance of extra systoles in an insured population *Circulation* 44 617 1971
 - 60 Blackburn H Taylor H L and Keys A The electrocardiogram in prediction of five-year coronary heart disease incidence among men aged forty through fifty nine *Circulation* 41(Suppl I) 154 1970
 - 61 Hinkle L E Jr Carver S T and Steve M The frequency of asymptomatic disturbances of cardiac rhythm and conduction in middle-aged men *Am J Cardiol* 24:49 1969
 - 62 Vedin J A Wilhelmsson C E Wilhelmsson L and Bjure J The relationship of resting and exercise induced ectopic beats to other ischemic manifestations and to coronary risk factors *Am J Cardiol* 30 75 1971
 - 63 Fulton M and Julian D G The Edinburgh chest pain study Geneva 1971 WHO working document Euro 8204 (3) pp 21 27
 - 64 Wilhelmsson L Elmfeldt D Tibblin G Vedin A and Wilhelmsson C Pre-coronary care Geneva 1972 WHO working document Euro 8204 (4)
 - 65 Norris R M Brandt P W T and Coughlin D E A new coronary prognostic index *Lancet* I 274 1969
 - 66 Elmfeldt D and Wilhelmsson L A study of representative post myocardial infarction patients aged 27 55 in Tibblin G Keys A and Werkö L editors *Preventive cardiology* Stockholm 1972 Almqvist & Wiksell J Wiley & Sons New York
 - 67 Vedin J A and Wilhelmsson C E Evaluation of a post myocardial infarction outpatient clinic in Tibblin G Keys A and Werkö L editors *Preventive cardiology* Stockholm 1972 Almqvist & Wiksell J Wiley & Sons New York
 - 68 Denborough M A Lovell R R H Nesher P J and Goble A J Arrhythmias and late sudden death after myocardial infarction *Lancet* I 386 1968
 - 69 The coronary drug project research group (In press 1972)
 - 70 Lown B Arrhythmias and sudden death Geneva 1971 WHO working document Euro 8204 (3) pp 17 19
 - 71 Lown B and Wolf M Approaches to sudden death from coronary heart disease *Circulation* 44 130 1971
 - 72 Kosowsky B D Lown B Whiting R and Guiney T Occurrence of ventricular arrhythmias with exercise as compared to monitoring *Circulation* 44 826 1971

with severe iron deficiency anemia. When severe myocardial hypoxia persists in such cases myocardial contractility becomes impaired and the severity of the heart failure increases.

Clinical features

The onset of symptoms in the infant who is developing congestive heart failure may be rapid and catastrophic as in patients with the hypoplastic left heart syndrome but often is quite insidious.

Feeding problems. An infant in congestive heart failure may tire after sucking for only a short while. He may accept a small amount of milk nap briefly from the exhaustion of sucking and then awaken, crying from hunger. This type of feeding history is almost invariably associated with a poor rate of weight gain. Failure to thrive is usual in infants with heart failure.

Tachypnea. Heart failure should be suspected when an infant's respiratory rate exceeds 60 per minute in the absence of pulmonary infection or metabolic acidosis since tachypnea is frequently the earliest sign of left heart failure in infancy. The respirations in such cases may be labored and are frequently associated with subcostal retractions. An expiratory grunt or wheezing may be present. Congestion of the bronchial mucosa may cause a chronic cough in some infants and may predispose them to repeated infections. Rales are usually heard late in the development of heart failure in infancy and their presence usually indicates that a severe degree of left heart failure exists.

Tachycardia. A cardiac rate in excess of 150 per minute is almost always present among infants with congestive heart failure. A rate in excess of 200 per minute may develop in these patients. However in the presence of such a rate one must distinguish between the sinus tachycardia of heart failure and paroxysmal supraventricular tachycardia arising from an ectopic focus. In early infancy the latter usually causes a fixed cardiac rate in excess of 250 beats per minute.

Cardiac enlargement. Cardiomegaly is virtually a *sine qua non* for the diagnosis of congestive heart failure in infancy and the configuration of the enlarged heart is frequently of assistance in diagnosing the

underlying cardiac defect. Cardiac failure in the absence of cardiomegaly is rare but may be seen when there is infradiaphragmatic insertion of anomalous pulmonary veins into the inferior vena cava or in some cases of cor pulmonale. In these instances radiographic changes in the appearance of the lung fields are usually of assistance in making the diagnosis.

Hepatomegaly. The liver is enlarged and readily palpable in infants with right heart failure. When failure develops rapidly the liver edge is rounded and may be quite tender from capsular distension; however the liver may be quite hard if the congestion is chronic. It is difficult to evaluate the significance of a palpable liver in an infant with pulmonary hyperaeration (e.g. in the presence of bronchiolitis) since the liver may be displaced downward in such cases by the depressed diaphragm but be normal in size.

Sweating and pallor. These may be prominent clinical signs in some cases. They result from the increased secretion of catecholamines that occurs as the body attempts to maintain compensation.

Cyanosis. If a patient has cyanotic congenital heart disease (a right to left shunt) e.g. transposition of the great arteries the cyanosis usually becomes more severe with the onset of heart failure. Other patients may develop cyanosis during heart failure due to either decreased oxygen diffusion at the alveolar level or to increased extraction of oxygen at the capillary level.

It is not uncommon to see either right or left heart failure in pure form in infants who have congenital heart disease. Thus pure left heart failure may occur in cases of coarctation of the aorta and aortic stenosis and marked tachypnea may be the initial sign of cardiac decompensation in such patients. On the other hand pure right-sided failure with hepatomegaly as the predominant sign may be encountered in the presence of tricuspid valve disease or pulmonic stenosis. More commonly however signs of left and right heart failure occur together since a reduction in left ventricular output occurs eventually in all cases. This reduced output leads to a diminished renal plasma flow, a rise in aldosterone secretion, progressive sodium and fluid retention, excessive weight gain

Table I *Leading causes of heart failure in infancy*

| | |
|---|---|
| I Congenital cardiovascular malformations | |
| A Large left to-right shunts | |
| 1 | Ventricular septal defect |
| 2 | Patent ductus arteriosus |
| 3 | Atrioventricular canal |
| 4 | Total anomalous pulmonary venous return |
| 5 | Truncus arteriosus with increased pulmonary vascular flow |
| 6 | Aortic pulmonary fenestration |
| 7 | A V fistula (especially cerebral) |
| B Large right to-left shunts | |
| 1 | Complete transposition of great arteries |
| 2 | Tricuspid atresia |
| 3 | Hypoplastic left heart syndrome with patent ductus arteriosus |
| C Obstructive lesions | |
| 1 | Coarctation of aorta (usually associated with patency of the ductus arteriosus) |
| 2 | Aortic stenosis |
| 3 | Pulmonic stenosis |
| 4 | Mitral stenosis |
| II Endomyocardial diseases | |
| A Endocardial fibroelastosis | |
| B Myocarditis (especially Coxsackie B) | |
| C Cardiac form of glycogen storage disease | |
| D Ectopic origin of left coronary artery from pulmonary artery | |
| E Medial necrosis of coronary arteries | |
| III Arrhythmias | |
| A Paroxysmal atrial or ventricular tachycardia | |
| B Paroxysmal atrial flutter | |
| IV Circulatory congestion | |
| A Overtransfusion or overhydration | |
| B Renal insufficiency | |
| C Anemia | |

secretion of norepinephrine which has a direct inotropic effect on myocardial contractility and which increases both systemic vascular resistance and venous tone. When such compensatory measures become inadequate progressive clinical heart failure results. In infants with large left to right shunts, failure most often occurs during the first six months of life.

Right to left shunts (Group IB Table I) In the presence of large right to left shunts severe arterial desaturation results from venous blood shunting directly into the systemic circulation. Thus, in addition to volume overloading of one or both ven-

tricles, these lesions lead to severe hypoxemia which impairs myocardial perfusion and which may lead to severe metabolic acidosis. The acidosis impairs myocardial contractility further and profound heart failure may occur.

Obstructive lesions (Group IC, Table I) These impose an impedance to ejection (systolic overload). The initial compensatory response in such cases is myocardial hypertrophy rather than dilatation. The resultant hypertrophic myocardium requires an increase in total coronary arterial flow. However, there is a greater impairment of flow in subendocardial vessels due to the high intramural pressure and coronary perfusion may become inadequate for the hypertrophic myocardium. In such cases depressed S-T segments ("strain pattern") become evident in the precordial leads and congestive heart failure or fatal dysrhythmias may follow.

Endomyocardial diseases (Group II Table I) These diseases depress myocardial contractility directly. End diastolic volumes increase as the ventricular ejection force diminishes and both ventricular dilatation and hypertrophy may become extreme.

Arrhythmias (Group III Table I) When persistent for more than 24 hours arrhythmias cause congestive heart failure in the majority of infants affected. The rapid rate of ventricular contraction causes an increase in the coronary blood flow needed to sustain the myocardium. However the short duration of diastole permits little time for coronary filling and myocardial ischemia eventually occurs. If there is an underlying structural abnormality heart failure occurs sooner after the onset of paroxysmal tachyarrhythmia than when the heart is otherwise normal.

Circulatory congestion (Group IV Table I) When there is an excessive intravascular volume (diastolic overload) due either to overhydration or severe renal insufficiency signs and symptoms of congestive heart failure may appear even though the myocardium is ejecting a larger than normal blood volume (high output failure Group IV). The use of diuretics to reduce the distended venous circulation may be more effective than the administration of cardiotonic agents in such a situation. High output failure may also be seen in association

young infants. In this instance the immaturity of their renal function places young infants into a similar category with any patient with renal insufficiency.

The activity of the enzyme β glucuronidase in the small bowel of young infants is relatively higher than it is in the intestinal tract of older individuals. Since many drugs and other potential toxic substances are excreted in the bile as water soluble glucuronide conjugates, the parent compounds (or one of their toxic metabolites) are regenerated when their metabolites undergo hydrolysis catalyzed by this enzyme. The resulting compounds are usually quite lipid soluble and are reabsorbed into the systemic circulation. This phenomenon of enterohepatic recirculation of compounds that are metabolized to glucuronides renders young infants less able to excrete many potentially toxic substances than older children.

It is impossible to apply our current knowledge of the pharmacology of most drugs in such a way that we can predict the immediate or delayed toxic effects of each agent in young infants. Thus we have no notion of the effect of various drugs upon the production or distribution of hormones whose actions in the first few weeks of life may be especially critical for the development of an infant's normal intellectual capacity such as thyroxine and insulin. Furthermore we are not certain of the safety of barbiturates in the neonatal period since these drugs are highly concentrated within the central nervous system

of the young of some laboratory species and remain there for prolonged periods. There are many other questions that can be asked about drugs that are used in the newborn period for which no dependable answer can be given. However it is possible to avoid the use of drugs that are known to displace bilirubin from its extracellular binding sites when treating young infants. It is also possible to exercise care when using agents that may build up within the infant because of his metabolic immaturity and others whose excretion may be slowed by his renal immaturity. Certainly physicians who must use pharmacological agents in newborn and other young infants have a right to expect reasonable answers to questions concerning the pharmacology of each new agent in such patients as the drugs are introduced. The rapid development of the study of the pharmacology of drugs in the immature and young of many species is new and it will require some time before this expectation can be realized.

REFERENCES

- Braunwald E. The control of ventricular function in man. *Br Heart J* 27:1 1965.
- Mitchell S C, Korones S B and Berenice H W. Congenital heart disease in 56 109 births. Incidence and natural history. *Circulation* 43:323 1971.
- Talner N S. Congestive heart failure in Moss and Adams editors. Heart disease in infants, children and adolescents. Baltimore 1968. The Williams & Wilkins Company pp 1004-1077.
- Yaffe S J, editor. Pediatric pharmacology. *Pediatr Clin North Am* 19:1 1972.

and occasionally peritoneal and peripheral edema

Infants with congestive heart failure are at great risk of succumbing rapidly to their disease or to intercurrent infection. The prompt institution of therapy to correct their physiological abnormalities is mandatory if they are to survive long enough for a palliative or corrective surgical procedure to be performed. If appropriate steps are taken to treat his congestive heart failure even a small premature infant with serious cardiac defects can be salvaged.

Concepts in developmental pharmacology

Drugs used to treat patients with cardiovascular disorders alter physiological and chemical processes by interacting at a molecular level with receptors in the various tissues or organs. Thus the activity of such drugs is dependent basically upon the nature and number of its effector sites and upon the physicochemical characteristics of the drug itself. Since effector sites probably have similar characteristics in most individuals their number and the factors that control the moment to moment concentration of the drug in the body are the important determinants of drug potency and safety. The number of effector sites in any given tissue or organism cannot be determined at the present time.

The concentration of a drug in the body at any point in time is determined by a complex set of kinetic constraints that describe its absorption, distribution, metabolism, and excretion. These constraints vary not only with the nature of the drug and its route of administration but also with numerous intrinsic host factors. Thus host factors play a major role in controlling the net amount of drug in the body at any time. These factors include the patient's size, the other drugs he is taking, the nature of his disease, his genetic endowment, and his physiological maturity.

The current interest in developmental pharmacology is the direct result of untoward occurrences that followed the use of drugs in young infants without regard for the added hazards to them imposed by immaturity of the systems that control drug distribution, metabolism, and excretion. Some of these are summarized below.

Distribution. All newborn infants develop some degree of hyperbilirubinemia during the first several days of life. The distribution of bilirubin in the body is controlled to a great extent by the number of binding sites available for it on the plasma albumin. In most cases, the distribution of the pigment is such that the cells of the central nervous system are not affected by its cytotoxic action. However, when the albumin is not capable of maintaining this favorable equilibrium, serious brain damage and even death may result. Thus the use of such bilirubin displacing drugs as the antibacterial sulfonamides, the sulfonamide diuretics, and strong acids (e.g., ethacrynic acid) is contraindicated during the hyperbilirubinemic stages of early life.

Metabolism. Most of the enzymatic reactions that convert drugs into their various metabolites in the body are catalyzed at a slow rate in young infants. Agents that require extensive metabolic alteration before they are eliminated must therefore be given to such patients in adjusted doses. Failure to recognize this can result in the accumulation of active drug within the body and can lead to serious toxicity. When chloramphenicol was administered to newborn infants in usual doses (adjusted only for the infant's weight), many infants developed the gray syndrome and died. It is now known that this syndrome of abdominal distension, respiratory distress, and circulatory collapse associated with a peculiar grayish color to the skin is related to the toxic effects of extremely high concentrations of the active drug in the body. It occurred in young infants because chloramphenicol is an antibacterial agent which requires metabolic alteration before it can be excreted and which built up in the infants due to their inability to convert it to its less toxic excretable metabolites. The "gray syndrome" was avoided in later years by adjusting the dose of chloramphenicol for the infant's metabolic immaturity as well as for his size and weight.

Excretion. Renal blood flow, glomerular filtration, and tubular function are all diminished in young infants. Accordingly, drugs that are excreted unchanged in the urine, or those whose water-soluble metabolites may be toxic in high concentration, must be administered with great care to all

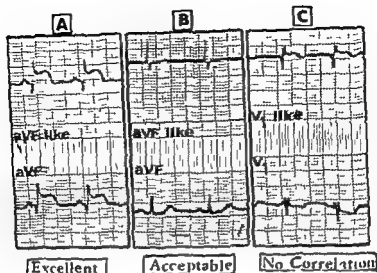


Fig. 1 Examples of the degree of correlation in simultaneously recorded conventional lead (lower tracing) and monitoring lead (upper tracing) ECG. See text for explanation.

and shape of QRS and P waves though the complexes were not identical. No correlation describes no similarity in form and/or direction (Fig. 1). In the V_1 -like lead, excellent correlation was found in 7 patients (86.6 per cent), acceptable correlation in 7 (11.6 per cent) and no correlation in only 1 case (1.7 per cent). Excellent correlation was present in 49 patients (51.6 per cent) with the aV_F -like lead, acceptable correlation in 9 (15 per cent) and no correlation in 2 (3.4 per cent). In the cases in which no correlation was found the QRS complex in the conventional lead was always isodiphasic thus probably explaining why small changes in the electrode position resulted in significant differences in the shape of the complexes recorded by the monitoring lead.

The almost insignificant number of patients in whom lack of correlation was found when comparing monitoring and conventional leads confirms the postulate that the basic diagnostic conclusions derived from the conventional ECG can be safely and reliably applied to the features obtained in a properly placed bipolar chest lead.

In setting up a continuous ECG monitoring system the adoption of a standard method of electrode placement seems advisable since such a method

allows easier interpretation of ECG data by the unit's staff and provides adequate information for more sophisticated computerized systems of arrhythmia detection as well.

N. Cristal M.D.
Coronary Care Unit
Negev Central Hospital
Beer Sheva, Israel

I want to thank Mrs. R. Hoffman R.N. for technical assistance in the preparation of this work.

REFERENCES

1. Stock J P F. New frontiers in arrhythmias. *Br Heart J* 33:809 1971.
2. Desautels R W, Block P and Hutter A M. Tachyarrhythmias in myocardial infarction. *Circulation* 45:681 1972.
3. Marriott H J L and Fogg E. Constant monitoring for cardiac dysrhythmias and blocks. *Mod Concepts Cardiovasc Dis* 39:103 1970.
4. Cristal M, Gueron M and Hoffman R. V_1 -like and aV_F -like leads for continuous electrocardiographic monitoring. *Br Heart J* 34:696 197.

Natural history of childhood lipid nephrosis

Recent studies have shown that the vast majority of children with the nephrotic syndrome have minimal histological changes by renal biopsy. This type of renal disease variously termed "nil disease

epithelial cell disease and lipid nephrosis" is characterized by complete remissions induced by glucocorticoid therapy. Sixty-one children with the nephrotic syndrome defined by heavy proteinuria, hy

Annotations

A sign of cardiac arrest

The physician is frequently confronted with a patient who complains of suddenly losing consciousness. When seen by his physician the patient fails to display signs or symptoms which make it possible to establish the cause. As a result the diagnosis is not established and management is unsatisfactory. Such patients are often considered to have a Stokes-Adams attack whereas other patients are told that they have experienced a simple fainting or syncopal episode. However, careful history taking from the patient and from anyone who has observed him faint can provide important and almost diagnostic information. The patient who suddenly develops unconsciousness due to cardiac arrest often experiences a generalized flushing and warmth or burning and tingling sensation of the skin of the entire body. He develops a marked redness or blush due to vaso-

dilatation of the veins and other vessels of the skin and then suddenly falls unconscious. Careful questioning of the patient when he recovers from cardiac arrest or Stokes-Adams attacks will often reveal this fairly characteristic syndrome. On the other hand patients with the common forms of syncopal episodes gradually or relatively slowly become extremely pale to ashen gray, sweaty, weak, giddy, and then faint. They do not display the manifestations noted above for sudden cardiac arrest.

George E. Burch, M.D.
Department of Medicine
Tulane University School of Medicine
1430 Tulane Ave.
New Orleans, La. 70112

Standard monitor leads— As reliable as conventional leads?

Two leading articles in the field of cardiac arrhythmias^{1,2} have recently re-emphasized the need for a uniform lead system in continuous electrocardiographic (ECG) monitoring. Doctors and nurses trained in the interpretation of the conventional ECG are often unable to make a diagnosis when dealing with an arbitrarily placed bipolar chest lead and immense quantities of ECG information are overlooked in thousands of patients being monitored daily. In addition, the introduction of more automated methods of arrhythmia detection require standard ECG data which can feasibly be converted into simple computer programs.

In an attempt to overcome the limitations of arbitrarily placed electrode, Marriott and Fogg³ described a set position which permits recording of a lead closely resembling the conventional V₁ lead. A modification of this method was suggested⁴ which likewise produces a V₁ pattern that can be easily changed into an aV_r pattern without altering the position of the chest electrodes on the patient.

We believe that more and more Coronary and Intensive Care Units will adopt one of these systems and comparable standard tracings will then be available in an increasing number of patients.

The question arises to what extent do the electrocardiographic features of the monitoring leads conserve the diagnostic implications of the conventional ECG leads.

We tried to find the answer to this question by recording parallel data in 60 consecutive unselected patients and evaluating the degree of correlation between the monitoring and the conventional leads. The conventional and the monitoring leads were simultaneously recorded on a two-channel recorder (Sanborn 7212) equipped with 2 ECG pre-amplifiers. Chest electrodes were placed as previously described.⁴ Briefly, electrode I was placed at the usual V₁ position, electrode II was placed below the outer third of the left clavicle and electrode III was placed at the left anterior axillary line over the ninth to tenth intercostal space. When electrode I is positive, II negative and III ground, a V₁-like lead is recorded. If electrode I is connected to the ground wire and III to the positive pole, an aV_r-like lead is recorded.

The results were tabulated as follows: an excellent correlation describes the situation where all the details of the ECG were identical. An acceptable correlation refers to a similarity in direction

three all had persistent proteinuria two had or had elevated BUN (20 and 30 mg per 100 ml respectively) and serum creatinine (2.7 and 3.0 mg per 100 ml respectively) and the remaining patient had an elevated blood pressure (150/100 mm. Hg). Four patients had died three of these were steroid resistant and had renal insufficiency at the time of death the fourth was steroid responsive and died of fluid and electrolyte complications during a relapse.

These data indicate that the nephrotic syndrome in preschool age children is not a benign disease despite the high responsiveness to treatment and the nature of the histologic lesion. Although the incidence of progression to renal failure remains low and for most children there is a marked reduction in the frequency of relapses by the time they reach

adolescence recurrences will be experienced by the majority of patients for many years.

Norman J Siegel MD
Assistant Professor of Pediatrics
Yale University School of Medicine
333 Cedar St
New Haven Conn 06510

REFERENCES

- 1 White R, H R, Glusko F F and Mills R J. Clinicopathological study of nephrotic syndrome in childhood. *Lancet* 1:1333 1970.
- 2 Siegel N J, Goldberg H, Krassner L S and Hayslett J P. Long term follow up of children with steroid responsive nephrotic syndrome. *J Pediatr* 81:251 1972.

The distress of dying

In contrast to many subjective reports few objective assessments have been made of the distress of dying. Doctors spend little time with dying patients so the observations recorded by nurses can be of particular value in providing a more comprehensive assessment than most doctors make alone. For this reason charts were devised that enable nurses to record the distress of dying in a manner similar to their recording of temperature (TPR) charts.

The factors to be assessed include pain, respiratory distress at rest, awareness of dying, depression, anxiety, vomiting and incontinence. Pain is recorded on a vertical chart graded 0 to 10 with the absence of pain recorded as 0 and very severe pain as 10. A similar principle is used for recording the other features of distress. The chart provides a pictorial image the significance of which is easily assessed. In extracting data for statistical analysis one has the problem of taking a reading from a vertical linear chart. Information recorded in a manner similar to that normally used for temperature chart. One can take the highest, lowest or average figures though where the aim is to explore distress the highest figures are most important and these are the ones used.

Data was obtained about 50 consecutive deaths occurring at home or hospital during a 12 month period in the Llandilo area of Wales. Significant differences occurred between deaths at home and in hospital for three reasons. Patients dying at home were (1) more likely to be fully alert shortly before death ($P < 0.05$) (2) less likely to be suffering from vomiting, incontinence or bed sores ($P < 0.001$) and (3) less likely to have unrelieved physical distress ($P < 0.05$).

It is possible that the presence of an alert mind and the absence of symptoms requiring constant

nursing care determined the site of death of these patients. Most people dying at home were free from vomiting, incontinence and bed sores whereas 28 per cent of all patients were troubled at some time by vomiting, 30 per cent had bed sores, 30 per cent had fecal incontinence, 2 per cent a colostomy, 20 per cent required an indwelling catheter and another 46 per cent had urinary incontinence. Only 26 per cent had none of the above symptoms.

Twenty two per cent revealed an awareness that they might die, 10 per cent expected to die soon, 10 per cent thought they would probably die, and 2 per cent thought they might possibly die. Fifty four per cent experienced pain of varying severity and 26 per cent had severe or very severe pain. Only 20 per cent had no respiratory distress at rest, 26 per cent had severe or very severe respiratory distress though in some of these instances the respiratory distress was mitigated by the fact that they were stuporose or comatose. Fifty six per cent experienced some anxiety and 40 per cent had depression. At times the anxiety of 14 per cent was so severe they could be described as fearful.

The results can be compared with those reported by Hinton¹ and Exton Smith² though the three surveys do differ in method, type of patient and area of environment. In Llandilo consecutive deaths in a community were assessed by trained nurses. The 220 patients in Exton Smith's survey all died in a geriatric unit. His information was obtained from personal observation and close enquiry from the nursing staff. Hinton's study was confined to a teaching hospital. He interviewed 102 dying patients and later questioned the sister or staff nurse about the state of the patient in the last few hours of life. On average the oldest patients were in Exton Smith's group² and the youngest in Hinton's¹.

The cause of death differed in the three surveys.

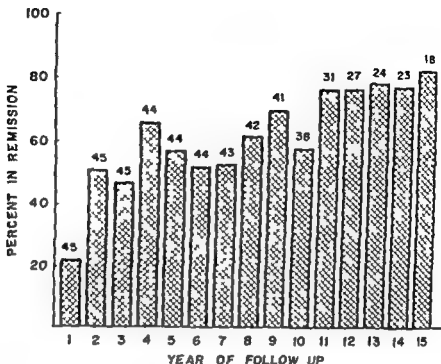


Fig. 1 The percentage of the relapsing group in complete remission during each year of follow up is shown by the bar and the number of patients remaining in the group is indicated at the top of each bar

poalbuminemia and edema and who had no known etiology for their disease formed the basis of our report.² Clinical criteria used in selecting these patients were: minimal follow up period of 5 years; age of onset 12 to 72 months; and complete remission due to initial steroid therapy. White and colleagues¹ have demonstrated that these clinical characteristics will define a homogenous group of patients with lipid nephrosis. Follow up was 10 years for 51 patients, 15 years for 21 patients and the average period of observation was 13.7 years.

There were 10 patients who experienced only the initial episode and 51 patients who had one or more recurrences. There were 47 boys and 14 girls and the mean age of onset was 37 months. Clinical features during the initial episode such as hematuria, transient hypertension and azotemia were not helpful in predicting whether an individual patient would follow a relapsing or non relapsing course. The most helpful feature in determining the likelihood of a relapsing course was the duration of remission after the initial episode. Twenty-two patients were without relapse during the first year after onset and of these 12 (55 per cent) subsequently had at least one relapse. Of the 13 patients who were in remission during the first two consecutive years from onset, 23 per cent subsequently relapsed while only one of eleven patients without relapse during the first three consecutive years had a later recurrence. All 10 patients who did not relapse during the first four consecutive years after onset remained in remission until the end of follow up. These data suggest that the relapsing form of lipid nephrosis is evident during the first two years of the disease and that complete resolution or cure is likely in the absence of a recurrence during the first three

years after onset. In contrast to patients with a non relapsing course, once a patient had experienced even one relapse, the likelihood of predicting complete resolution even after extended periods in remission was uncertain. For example, of 31 patients in remission for five years at some time during the course of illness after a relapse, seven (22 per cent) subsequently had a recurrence.

Sufficient data were available on 45 of the 51 cases with a relapsing course for long term evaluation of the frequency and pattern of relapse. While only 23 per cent remained free of relapse during the first year, approximately one half of the group was in remission during each of the subsequent six years. Although the proportion in remission gradually increased with time, 20 per cent of the group had relapses as long as 15 years after onset (Fig. 1). Similarly, during the first year after onset the group averaged more than one relapse per patient but thereafter the relapse rate was less and a distinct trend toward a decreasing rate was evident after the tenth year of disease.

The clinical status of all 61 patients at the end of the study was evaluated by complete history, physical examination, urinalysis, blood urea nitrogen (BUN) and serum creatinine. The ten patients who had only the initial episode each had a normal BUN, serum creatinine and blood pressure at the completion of follow up. Of the 51 patients with a relapsing course, 25 had been in remission for a minimum of two consecutive years, 22 were continuing to experience relapses, 13 of these were steroid responsive, 6 were steroid dependent and 3 were steroid resistant. Of these 47 patients, all had normal BUN, serum creatinine and blood pressure except the three who were steroid resistant. Of these

Letters to the Editor

Lidocaine in ventricular arrhythmia

To the Editor

Lidocaine is considered to be the drug of choice in the initial treatment of ventricular arrhythmias especially during an acute myocardial infarction. Electrophysiologic studies by Rosen and co-workers¹ using a single bolus of lidocaine and Josephson and co-workers² using a bolus of lidocaine followed by an intravenous drip of 1 mg. per minute demonstrated no significant effect on A-V nodal or His-Purkinje conduction time in man. The recent article by Josephson and colleagues on the effects of lidocaine on refractory period in man³ observed the development of complete A-V nodal block with ventricular asystole in one patient followed by persistent delay in His-Purkinje system conduction upon return to normal sinus rhythm.³ We have recently observed a similar patient who developed asystole and complete heart block following a bolus injection of lidocaine.

A 49-year-old Caucasian male physician weighing 140 lbs. sustained an anterior wall infarction complicated by a right bundle branch block, left anterior hemiblock, myocardial aneurysm and congestive heart failure in December 1971. On December 9, 1972, he was admitted to the CCU with a chief complaint of severe crushing substernal chest pain of several hour duration. His medications consisted of digoxin 0.25 mg. and Easidrix 50 mg. daily. His initial serum K^+ was 4.1 mEq. Because of frequent premature ventricular contraction and short runs of ventricular tachycardia, 80 mg. of lidocaine was rapidly administered as a bolus intravenously. Within 2 minutes, complete heart block with ventricular asystole developed; atrial activity, however, was not affected. Immediate closed chest massage and ventilation were initiated and ventricular escape beats emerged (Fig. 1). Within 10 minutes, regular sinus rhythm returned. A temporary pacemaker was then inserted to allow the use of continued antiarrhythmic medication—lidocaine. The patient died on the third hospital day following the development of refractory ventricular arrhythmias.

This experience would seem to be the second

documented episode of ventricular asystole following the administration of lidocaine. We have previously reported the occurrence of sinoatrial arrest as a complication of lidocaine.⁴ In the present case, lidocaine may have depressed conduction through the posterior division of the left bundle. We concur with the conclusions of Josephson and colleagues³ that smaller than average doses of lidocaine be administered to patients with congestive heart failure or liver disease of any etiology. In the presence of bilateral bundle branch block, lidocaine should probably be administered with caution and slowly.

Joseph P. Liss Jr. MD
Robert W. Jeresaty MD
Jadhour Akkhour MD
Department of Medicine
Section of Cardiology
Saint Francis Hospital
Hartford, Conn. 06105

REFERENCES

1. Rosen H, M. Lau S, H. Weiss M, H. and Damato A. N. The effect of lidocaine on atrio-ventricular conduction in man. *Am J Cardiol* 25:1, 1970.
2. Josephson M, E. Caracta A, R. Lau S, H. Gallagher J, J. and Damato A. N. Effects of lidocaine on refractory periods in man. *Am Heart J* 84:778, 1972.
3. Jeresaty R, M. Kahn A, H. and Landry A. B. Jr. Sinoatrial arrest due to lidocaine in a patient receiving quinidine. *Chest* 61:683, 1972.

Atrial dissociation

To the Editor

In the latest case report of Atrial Dissociation by Drs. Svendsen and Jorgensen (*Am Heart J* 80:103, 1973), it appears that an entirely different interpretation can be made of the electrocardiogram, the intracavitary electrogram and the alleged atrial activity.

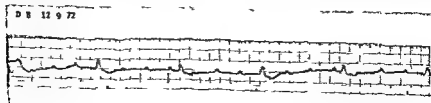


Fig. 1 Monitor strip showing complete heart block. Ventricular rhythm = 37 per minute, atrial rhythm = 125 per minute.

mainly because 100 per cent of Hinton's patients died from cancer. He reported that di comfort was not greatest in patients dying from cancer but patients dying from heart failure or both had most physical distress.¹

Hinton also pointed out that awareness of dying increases as death draws closer. 75 per cent of his patients indicated such awareness during the final weeks. The Llandiloos figure of 22 per cent is much lower, probably because the nurses did not seek this information but recorded only information volunteered by patients. It is similar to the 25 per cent reported by Exton Smith² so from these two studies it seems likely that about a quarter of dying patients indicate awareness of approaching death to their attendant nurses. Three and two-tenths per cent of Exton Smith's patients² and 10 per cent of those in Llandiloos expected to die soon while the figures given by Hinton¹ range from 6 per cent at the first interview to 20 per cent during the last week of life.

Impairment of consciousness increases as death approaches. 34 per cent of Hinton's patients¹ were unconscious for at least 24 hours before death and 40 per cent of those dying in Llandiloos were in the same state.

Thirteen and six tenths per cent of Exton Smith's patients² experienced moderate or severe pain. Hinton¹ found that 65.7 per cent had pain though the severity was not classified and 18 per cent of those with pain failed to receive adequate relief. In Llandiloos 54 per cent had pain with 26 per cent experiencing severe or very severe pain. Hinton¹ stated that dyspnea is more difficult to relieve than pain, nausea or vomiting and that out of 17 patients suffering from respiratory distress, 14 (82 per cent) failed to obtain relief.

A few patients (3.2 per cent) in Exton Smith's study² showed severe anxiety and were afraid to be left alone. Hinton¹ found that 64.7 per cent showed at least slight anxiety and 12.7 per cent experienced moderate anxiety but none were classified as fearful. In Llandiloos 56 per cent showed at least slight anxiety while at times 14 per cent were fearful. Hinton¹ included as severely anxious or fearful only patients who had been fearful, agitated or panic-stricken for most of the week. In Llandiloos any transient state of fearfulness that the nurse noted was included. Depression occurred in 7.7 per cent of Exton Smith's patients² while Hinton¹ found that 66.7 per cent experienced some depression and that 16.7 per cent experienced moderate or severe depression. In Llandiloos 40 per cent experienced at least slight depression while 26 per cent were moderately or severely depressed. Altogether 12 per cent of the Llandiloos patients wanted to die which is higher than the 5 per cent reported by Exton Smith.² Hinton¹ pointed out that 65 per cent of depressed and 52 per cent of anxious patients remain unrelieved.

II Dees Rees M D MRCGP
Abernant Llandiloos
Montgomeryshire SL18 6AU
E. Wales

REFERENCES

- 1 Hinton J M. The physical and mental distress of the dying. *Q J Med* 32:1 1963.
- 2 Exton Smith A N. Terminal illness in the aged. *Lancet* 2:305 1961.
- 3 Rees W D. The distress of dying. *Br Med J* 3:105 1972.

by Dr Orsiccio, Dr Chien and Dr Mädlar. However, we do not agree with their interpretation of the electrocardiograms.

First, the waves which we interpreted as right atrial activation with A-V dissociation occurred at the rate of 105 per minute (paper speed 50 mm per second). These waves could clearly not be due to diaphragmatic electrical activity. The morphology in intracavitary lead from different parts of the right atrium in our opinion excludes any other possibility than that of a right atrial P wave.

Second, we are not convinced that our identification of the QRS complex is wrong. Recordings from the distal part of the right atrium close to the tricuspid valve which was not published in our paper strongly suggest that the largest waves seen in Lead V₁ are QRS complexes. Additional support to our interpretation is found in our present Fig. 1 which shows another strip of the electrocardiogram. A slightly irregular rhythm is seen with one premature

beat. The relation between what we interpret as the P wave and the QRS complex is constant except in the beat following the premature beat when the PR interval is slightly shortened. However, the correct identification of the P and the QRS complexes is not essential for the diagnosis of atrial dissociation if our interpretation of the extra right atrial P wave is accepted.

Discussing the interpretation of the electrocardiogram yet another alternative deserves mentioning, namely the possibility that both waves belong to the QRS complex—i.e. gross intraventricular conduction disturbance is present. The ECG should then be one with atrioventricular dissociation and intraventricular conduction defect. However, we do not feel that this is the correct interpretation.

Egil Sivertsen M.D.
Leif Jørgensen M.D.
Lillelø Hospital, Department VIII
Oslo, Norway

In Fig. 1 the initial PR interval is about 0.24 second. The ST segment is elevated in V_1 and V_2 . This appears to be an acute interseptal infarction with an old inferior infarction. The PR prolongation suggests severe septal damage. Notice appropriately the QRS is followed by a T wave.

In Fig. 2 what is identified as a QRS complex is not followed by a T wave. Therefore this is probably the P wave. What was identified as the P wave is the QRS complex. Notice the large S in LI with left axis deviation. It is now apparent that what was identified as a P wave in V_1 is found to be a QRS complex with an RSR pattern. The PR interval would now be about 0.54 second or more further prolonged over what was initially seen rather than being shortened as suggested by the authors' interpretation. There is little doubt that this is trifascicular block. The acute anterior wall infarction is demonstrated by the loss of precordial R wave with ST segment elevation. The T wave is lost in the P wave. The giant P waves suggest severe pulmonary hypertension. It is of interest to speculate that the elevated PR segment in Leads I, aVR, aVL and V_1 through V_6 with PR segment depression in Leads II, III, aVF, V_1 and V_2 may represent atrial hypertension and atrial infarction.

If one carefully measures what is taken as an extra P wave in the right precordial lead and the atrial electrogram (again the P wave and the QRS misidentified) it is at the rate of about 50 per min.

If one were to refer back to the nurse's notes, tachypnea at this rate would be described.

What one is seeing is diaphragmatic electrical activity that we have described¹ as a preterminal sign in severe obstructive lung disease or congenital failure when severe hypoxia is present.

It is not uncommon to mistake diaphragmatic action potentials for dissociated atrial activity. We hope that the review by the authors is in agreement with us.

Ralph G. Orsiccio, M.D.
Chief of Medicine
Chin Chen Chien, M.D.
Associate Medical Resident
Ernesto Wadarang, M.D.
Associate Medical Resident
Saint Elizabeth Hospital
225 Williamson St.
Elizabeth, N.J.

REFERENCE

1. Orsiccio W. Atrial dissociation (Letter to the Editor) *Am J Cardiol* 29:303 1972.

Reply

To the Editor

We appreciate very much the interest in our case report of atrial dissociation and the comments made

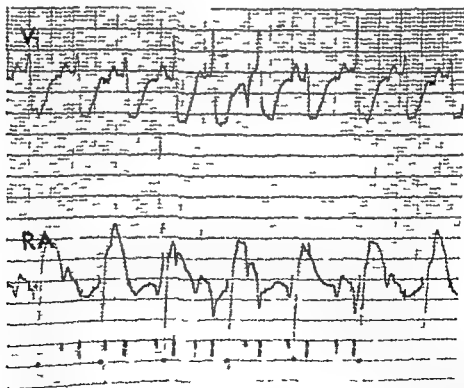


Fig. 1. Electrocardiogram showing Lead V_1 and intracavitary right atrial (RA) lead. A close time relationship is found between the waves which are interpreted as P waves (marked '1') and those interpreted as QRS complexes (marked '2')—but not in the opposite direction—in spite of irregular rhythm. Independent of these waves large negative complexes (marked '3') are seen in the RA lead, interpreted as right atrial P waves. Paper speed = 25 mm per second.

perform diagnostic wonders at the bedside. This book should interest all students, interns and residents as well as doctors in practice who wish to become proficient in examination of the heart. This is a valuable publication.

✓ **SPONTANEOUS HYPERTENSION: Its Pathogenesis and Complications**. Edited by Hozo Okamoto. M.D. Tokyo 1972. Igaku Shoin Ltd. 266 pp. Price \$74.80.

Dr Okamoto's book contains the papers presented at a seminar held in Kyoto from October 18 to 22, 1971, on a strain of rats that develop hypertension spontaneously. Investigators from many parts of the world who are studying various aspects of this syndrome in the rats participated in the conference. This publication consists of 8 chapters. Several papers are included in each chapter. The subjects discussed were development and heredity, catecholamine metabolism, neural factors and behavior, cardiovascular dynamics, cardiovascular pathology, endocrine factors, renal factors and electrolytes, and essential hypertension and spontaneous hypertension. These presentations of this interesting phenomenon extensively review the pathophysiology of an interesting disease and animal model of hypertension. Regardless of its relationship to hypertension in man, this book is an extremely important one. The seminar gathered together a great deal of valuable information for others who are interested in hypertension. This is an important publication.

✓ **CEREBRAL VASCULAR DISEASES: Eighth Conference**. Fletcher H. McDowell, M.D., Chairman, and Fletcher H. McDowell, M.D., and Robert W.

Brennan, M.D., Editors. New York 1973. Grune & Stratton, Inc. 319 pp. Price \$12.50.

The publication of the proceedings of the Eighth Conference on Cerebral Vascular Diseases reflects a meeting which resembles the preceding ones. This book contains discussions of extracranial arterial surgery and a preliminary report on the cooperative aneurysm project, various aspects of cerebral ischemia (pathology, microcirculation, metabolism, pathophysiology), head injuries, cerebral angiography, role of platelets and pharmacology. The papers are interesting and the appended discussions are even more important to readers who were not in attendance at the conference. This is another important publication which should interest all physicians, not only neurologists and neurosurgeons.

✓ **ESSENTIALS OF CARDIAC ARRHYTHMIAS: Diagnosis and Management**. Samuel Bellet, M.D. Philadelphia 1972. W.B. Saunders Company. 458 pp. Price \$15.50.

This book of 38 chapters on one of the most important problems in cardiology was written by Dr. Bellet shortly before he died of myocardial infarction. He had been interested in arrhythmias of the heart for many years and has written fairly extensively on the subject. As in previous writings, this book is intended for the practicing physician. It is well organized and clearly written. The 38 chapters cover all the common arrhythmias and their management as well as the anatomic and physiologic considerations necessary for an understanding of the arrhythmias. This is a well illustrated book and a good addition to the many others already available on the subject. Students, interns and residents will find the book to be useful.

Book reviews

✓ **ADVANCES IN MICROCIRCULATION** vol 4 H Harders editor Hamburg Basel New York 1972 S Karger AG 232 pp Price \$24.40

This fourth volume on Advances in Microcirculation should interest all physiologists and all physicians who treat cardiovascular diseases. It is divided into six parts related to microcirculatory blood flow, hemodilution and rheologic changes, effects of dietary fats, plethysmography, thermography, use of radionuclides, labelled colloids, and leukocyte sequestration in the pulmonary circulation in hemorrhagic shock. As are the past volumes, this is also a very good one. The reader will find this book to be interesting and useful; it is highly recommended.

✓ **RECENT ADVANCES IN RENAL PHYSIOLOGY** Editors H Wirtz and F Spinelli Basel 1972 S Karger AG 298 pp Price \$27.45

This publication contains papers presented at an international symposium held in Breitenberg, Switzerland, August 2 to 5, 1971. The sessions were concerned primarily with the handling of sodium by the kidneys. The many papers were related to mechanisms of sodium excretion, renal circulation, function of the medulla, and diversity of the nephrons. The contributors have studied water and electrolyte excretion rather extensively and gathered, as in all such symposia, to exchange views and findings. This gathering must have been interesting as reflected in the resultant publication. Those who are interested in the fundamentals of renal function will find this book to be stimulating and useful. The papers are thought-provoking in that they readily reveal the many questions yet to be answered. Renal physiologists will find this to be a valuable publication.

✓ **ANGINA PECTORIS** American Heart Association Monograph No. 37 Edited by Charles K. Friedberg M.D. with the assistance of Ephraim Donoso M.D. New York 1972 1227 pp Price \$5.00

The American Heart Association has for the past several years published symposia on various important cardiologic problems. This is the thirty-seventh on an extremely important clinical state which waxes and wanes in interest among clinicians. At present it is of considerable interest because of the surgical approaches to the management of coronary heart disease. This publication contains nothing new to cardiologists but it should interest others who fail to follow the medical literature closely. The subjects discussed are of routine interest though important—viz. coro-

nary angiography, bypass surgery, medical management, diagnosis. This monograph should interest students, young physicians, internists, and general practitioners.

NEURAL AND PSYCHOLOGICAL MECHANISMS IN CARDIOVASCULAR DISEASE Proceedings of a Symposium held in Stresa, Italy, July 19-21, 1971. Edited by Alberto Zanchetti. Milano 1972. Casa Editrice 387 pp Price \$25.00

These proceedings of an excellent symposium held in Stresa, Italy, July 19 through 21, 1971, on the psychological factors in cardiovascular disease should be welcomed by those who were unfortunate not to have been present at the sessions. Among the many subjects discussed were those related to methods for evaluating neural and psychological factors in cardiovascular disease, influence of personality and perception. The many interesting presentations were delivered by contributors from many parts of the world. The book is very good and reflects outstanding work of the organizing group from Milano. This is an important and interesting contribution to the medical literature. The series of papers should interest all physicians.

✓ **PHONOCARDIOLOGY** Integrated Study of Heart Sound and Murmurs. Jozef Wartak M.D. New York 1972 Harper & Row Publishers 156 pp Price \$17.50

The short monograph on heart sounds is a little different from the usual publications. The author presents the physics of sound in a simple fashion. He does this with good clear illustrations which the reader must study carefully. The relationship of heart sounds to other hemodynamic and to electrocardiographic phenomena is illustrated. These illustrations require thoughtful study and understanding. The normal and abnormal heart sounds and murmurs are well presented. The author refrains from discussion of controversial matters. This is a good idea for a book of this type. The book is useful for beginners.

✓ **AUSCULTATION OF THE HEART** 4th ed. By R. W. D. Turner. Edinburgh 1972 Churchill Livingstone 122 pp Price \$5.50

This small paperback on examination of the heart is not only an important and practical publication but a much needed one in this era of the cardiac catheter. A well-trained cardiologist can

Announcements

Appeal for rheumatic subcutaneous nodules

An appeal is made to physicians and investigators for rheumatic subcutaneous nodules recently formed in the course of rheumatic carditis for use in studies on the pathogenesis of rheumatic fever

Nodules situated on the extremities readily

accessible to biopsy and just in the process of forming before steroid therapy are especially so. It, although any nodules will be welcome. Self-addressed containers with fluids and details concerning biopsy technique may be obtained from Dr Bernice G Wedum c/o Mr Gun Hyuk Kim Bldg 37 Room 1C27 National Institutes of Health Bethesda Md, 20014

adverse actions can be terminated by stopping administration or rarely by the administration of propranolol

A more serious toxic effect has been described for isoproterenol myocardial infarction and necrosis. This occurs in experimental animals to which massive doses of isoproterenol have been given. The doses needed are at least 1 000 times larger than any therapeutic dose. And this effect is produced by any active catecholamine. This tissue damage is probably due to physical effects of intense myocardial stimulation, relative coronary insufficiency and direct biochemical effects of the catecholamines. There is no evidence that this adverse result can or has happened with any therapeutic or diagnostic use of isoproterenol.

Are there any drug interactions to worry about when using isoproterenol? Obviously *beta* blocking agents would interfere in the use of isoproterenol in bronchial asthma. However *beta* blockade is a property of a

fairly restricted chemical structure. Veratrum alkaloids would diminish the positive chronotropic action of isoproterenol. However drugs that interfere with adrenergic transmission such as reserpine, guanethidine, methylphenidate or cocaine would not affect the isoproterenol response. This catecholamine not being significantly taken up by the amine pump of the nerve end is not affected by denervation phenomena.

Isoproterenol should be the drug of choice whenever the effect of specific sympathetic nerve stimulation of the heart is desired. This conclusion is valid only at the present instant in time. There are indications that the cardiac *beta* receptors are different from smooth muscle *beta* receptors. Cardioselective *beta* blocking agents are undergoing clinical trial now. It won't be long until cardioselective *beta* agonists will be readily available. At that time our opinion of the use of isoproterenol may have to be changed.

terenol is unquestioned. There are only two questions: is a decongestant action needed also (by substituting epinephrine or adding phenylephrine), and how should the isoproterenol be administered? The latter question arises for any clinical use of isoproterenol. As a catecholamine its survival when administered orally is extremely variable. When first tested clinically in this department in the 1940's, isoproterenol was supplied as an oral tablet. This, however, was most unsuccessful. Some subjects showed no response; others showed extreme tachycardia and palpitations.

Three methods of administration are available: intravenous infusion for prompt, controllable cardiovascular effects; sublingual or buccal tablets for chronic systemic action; and inhalant solutions for local action in the bronchi.

The syncope of complete heart block is due to low cardiac output caused by episodes of either bradycardia or ventricular fibrillation. Unless the ECG is being continuously monitored it may be difficult to distinguish which condition is present. Isoproterenol is the drug of choice; it will correct bradycardia and there is reasonable evidence that it would not prolong a short episode of fibrillation. Transient complete heart block is readily treatable with isoproterenol. Permanent heart block is usually treated by pacemaker implantation.

The use of isoproterenol in cardiogenic shock is more uncertain. The hypotension and weak pulse associated with pulmonary embolism, open heart surgery, or acute myocardial infarction is presumed to be due to myocardial weakness. On this basis a rapid acting positive inotropic agent is indicated. This can be either isoproterenol or norepinephrine. Both agents affect the cardiac beta receptors to increase the force of contraction and improve cardiac output. Norepinephrine by adrenergic alpha receptor action can produce in addition vasoconstriction and a pressor response. Isoproterenol will tend to increase pulse rate; norepinephrine will tend reflexly to decrease pulse rate. Isoproterenol will tend to increase coronary flow, but by decreasing total peripheral resistance may adversely affect renal and cranial blood flow. Norepinephrine, while increasing coronary flow by beta receptor action and by increasing

arterial pressure by alpha action may be more seriously producing renal shutdown. And finally both drugs can produce serious arrhythmias. It should be noted that in experimental pharmacology an easy way to produce arrhythmia is simultaneous stimulation of the vagus and sympathetic nerve supply of the heart.

It seems obvious then that the choice of drug depends on the exact conditions in each patient being treated. This is shown by the clinical literature on the subject; success and failure for each drug; isoproterenol and norepinephrine is well documented.

In many ways isoproterenol should be of great value in cardiology. Administered intravenously its actions are predictable and controllable. Two examples of the experimental and diagnostic value of isoproterenol can be cited.

Many clinical studies of new and old adrenergic beta receptor blocking agents are being done at the present time. Any drug action purporting to be due to beta receptor blockade must be proved by comparison of the subject's response to isoproterenol before and after the experimental drug dosage. Isoproterenol activates all beta receptors so it can be used as a control for any beta blocking agent including the newer cardioselective agents such as practolol. Without this control other actions of the experimental drug may be producing the observed effect.

Isoproterenol increases the functional stenosis in idiopathic hypertrophic subaortic stenosis. This action has value during diagnostic and evaluative catheterization. Unfortunately the converse of this effect is not true: adrenergic beta receptor blocking agents are not often beneficial in this form of stenosis.

Is isoproterenol a safe drug? As with any drug, too much can be administered. In any reasonable therapeutic overdosage only two adverse things can happen: tachycardia and relative coronary insufficiency. Isoproterenol is less likely to produce severe arrhythmias than either epinephrine or norepinephrine. The coronary insufficiency is due to the disparity between the increased coronary flow and the even greater increased myocardial oxygen demand caused by the intense myocardial stimulation. The

adverse actions can be terminated by stopping administration or rarely by the administration of propranolol

A more serious toxic effect has been described for isoproterenol myocardial infarction and necrosis. This occurs in experimental animals to which massive doses of isoproterenol have been given. The doses needed are at least 1 000 times larger than any therapeutic dose. And this effect is produced by any active catecholamine. This tissue damage is probably due to physical effects of intense myocardial stimulation relative coronary insufficiency and direct biochemical effects of the catecholamines. There is no evidence that this adverse result can or has happened with any therapeutic or diagnostic use of isoproterenol.

Are there any drug interactions to worry about when using isoproterenol? Obviously *beta*-blocking agents would interfere in the use of isoproterenol in bronchial asthma. However *beta* blockade is a property of a

fairly restricted chemical structure. Veratrum alkaloids would diminish the positive chronotropic action of isoproterenol. However drugs that interfere with adrenergic transmission such as reserpine, guanethidine, methylphenidate or cocaine would not affect the isoproterenol response. This catecholamine not being significantly taken up by the amine pump of the nerve end is not affected by denervation phenomena.

Isoproterenol should be the drug of choice whenever the effect of specific sympathetic nerve stimulation of the heart is desired. This conclusion is valid only at the present instant in time. There are indications that the cardiac *beta* receptors are different from smooth muscle *beta* receptors. Cardioselective *beta* blocking agents are undergoing clinical trial now. It won't be long until cardioselective *beta* agonists will be readily available. At that time our opinion of the use of isoproterenol may have to be changed.

Changes in the coronary vasculature in endomyocardial fibrosis and their possible significance

Zilton A. Andrade M.D.*

Antonio R. L. Teixeira M.D.**
Bahia, Brazil

In the first study of the pathology of endomyocardial fibrosis that was published by this Department¹ a striking vascular lesion was described in one of the four cases presented. That lesion was interpreted as consisting of thrombosis with organization and recanalization. It involved medium sized and small arteries in the fibrotic endomyocardial scars. Other but less impressive changes such as medial fibrosis and intimal thickening, were seen in all the other cases. As further cases of endomyocardial fibrosis have come under microscopic observation there has emerged what in our opinion seems to be a peculiar pattern of vascular changes affecting the coronary vessels in this disease.

Although the gross and histological features of the heart itself have been thoroughly studied in cases of endomyocardial fibrosis,²⁻⁶ little attention has been paid to changes in the coronary vessels, specially those of smaller caliber. Moreover most reports refer to the presence of vascular sclerosis and occlusion within the densely scarred endomyocardial tissues. McKinney⁷

mentioned subendothelial deposits of eosinophilic material which decrease the vascular lumen. Connor and associates⁸ described an early lesion consisting of medial swelling due to the presence of increased amounts of amorphous polysaccharide material; they also noted a late change represented by vascular sclerosis and partial occlusion of the lumen.

As coronary arterial changes were observed to be frequent in our experience of 12 necropsies in cases of endomyocardial fibrosis we decided to make a systematic study and description of these changes and to try to interpret their nature and significance.

Material and methods

Complete necropsies were performed in 12 cases of endomyocardial fibrosis. Four of these cases (Cases 1, 2, 3, and 4) appeared in a previous report.¹ Brief general clinicopathological data are presented in Table I. Three of the patients presented with hepatosplenic schistosomiasis in all the others endomyocardial fibrosis was the

From the Department of Pathology Hospital Prof. Edgard Santos University of Bahia School of Medicine Salvador Bahia Brazil

Received for publication Sept. 12, 1972

Reprint requests to Dr. Zilton A. Andrade Department of Pathology Hospital Prof. Edgard Santos 40.009 Salvador Bahia Brazil

*Professor of Pathology University of Bahia School of Medicine

**Instructor in Pathology University of Bahia School of Medicine

Table 1 Clinicopathological data on 12 fatal cases of endomyocardial fibrosis

| No | Age (yr) | Sex | Race | Duration of illness | Heart weight (Gm) | Distribution of endomyocardial fibrosis | | Valvular involvement | Intracardiac thrombosis |
|----|----------|-----|------|---------------------|-------------------|---|-----|----------------------|-------------------------|
| | | | | | | R V | L V | | |
| 1 | 27 | M | N | 2 years | 400 | • | • | Mitral | Rt atrium |
| 2 | 24 | F | N | 7 years | 380 | 0 | 0 | None | None |
| 3 | 21 | M | N | 13 years | 280 | • | 0 | None | Rt atrium |
| 4 | 28 | F | N | 2 years | 350 | • | • | None | Rt atrium |
| 5 | 26 | F | N | 5 years | 670 | • | • | None | Rt atrium |
| 6 | 35 | F | N | 4 years | 400 | • | • | Mitral | None |
| 7 | 25 | M | N | 5 months | 450 | • | • | Tricuspid | None |
| 8 | 9 | F | N | 18 months | 200 | • | 0 | Tricuspid | Rt ventricle |
| 9 | 22 | M | N | 1 year | 400 | • | 0 | Tricuspid | Rt atrium and ventricle |
| 10 | 62 | F | M | 3 years | 350 | • | • | Mitral | Rt atrium |
| 11 | 21 | F | N | 8 months | 270 | • | • | None | None |
| 12 | 14 | M | N | 7 months | 180 | • | 0 | None | None |

Degree of endomyocardial fibrosis: 0 = none; 1 = mild; 2 = moderate; 3 = marked; 4 = severe. (•) = present; (0) = absent. (N) = Negro; (M) = Malay; (W) = White.

main pathological process. The hearts presented considerable endomyocardial scarring of one or both infarct tracts of the ventricles and in some instances also of the apex of one or both ventricles (see Table 1). The fibrotic areas were contracted, the other parts of the hearts showed dilatation. All the hearts were fixed in 10 per cent formalin. Tissue blocks were taken from several different areas in every case and embedded in paraffin. Sections five microns thick were stained by the following methods: hematoxylin-eosin, Mallory's triple chrome stain, periodic acid-Schiff (PAS), Weibert's iron-haematoxylin stain for collagen and elastic fibres, and Movat's pentachrome stain. Paraffin sections of practically every organ of the body were examined with particular attention to the localization of any vascular lesions. Chagas disease, a common condition in the area, is ruled out because no gross or microscopic evidence of that disease was ever present in the cases of endomyocardial fibrosis studied here.

Results

Vascular lesions were found mainly within the area of endomyocardial fibrosis which appeared as broad septa penetrating the myocardium from the densely scarred

endocardium. They were sometimes observed within the uninvolved myocardium and in the adipose tissue of the epicardium (Fig. 3) but no comparable changes were found in blood vessels outside the heart. Only coronary arteries and arterioles were involved. The coronary veins showed no more than edema of the media, congestion and occasionally thrombosis. For purposes of description the findings will be considered under the following headings.

Medial degeneration. Medial swelling was frequently seen. Sometimes a basophilic amorphous material separates the muscle cells; this material was PAS positive and showed the staining characteristics of amorphous polysaccharide material with Movat's pentachrome stain. There was swelling of the muscle cell cytoplasm with or without vacuolation. These medial changes were particularly severe at the sites of vascular bifurcation where foci of hyaline necrosis were also seen (Fig. 1).

Medial focal proliferation. Focal areas of muscular hyperplasia were seen in some medium-sized arteries, usually associated with some degree of medial degeneration or fibrosis.

Intimal proliferation. This appeared to be associated with deposition of fibrinous material on the endothelial surface of



Fig 1 A through F Several types (probably evolutionary stages) of medial degeneration affecting the coronary arteries in endomyocardial fibrosis. A Mucoid degeneration. The muscle cells in the media appear dissociated (Hematoxylin and eosin. Original magnification $\times 150$). B Vacuolation of the muscle cell cytoplasm (Hematoxylin and eosin. Original magnification $\times 130$). C Formation of clefts in the degenerated media (Hematoxylin and eosin. Original magnification $\times 130$). D Medial hypertrophy and sclerosis (Movat's pentachrome stain. Original magnification $\times 100$). E Medium sized subepicardial coronary artery showing medial fibrosis affecting only part of the vessel (Movat's pentachrome stain. Original magnification $\times 100$). F Medial degeneration with lumen not altered (Movat's stain. Original magnification $\times 130$).



Fig. 2 *A through I* Several stages of the lesions affecting the coronary arteries in endomyocardial fibrosis. *A, B, C* and *D* Small arteries showing intimal fibrin deposit on endothelial cell proliferation, intimal fibrosis and the formation of fibrous bands that divide the lumen. In *D* the media of the smaller vessel is almost completely replaced by fibrosis (Movat's pentachrome stain. Original magnification $\times 130$). *E* and *F* Intimal proliferation and the formation of secondary lumina. External to the endothelial cells there is a thin layer of fibrous material (Movat's pentachrome stain. Original magnification $\times 130$). *G, H* and *I* Multiluminal vessels. The medial coat is almost totally replaced by fibrosis. Probably an end-stage lesion (Hematoxylin and eosin. Original magnification $\times 130$).



Fig 3 An advanced change represented by medial fibrosis and the formation of new vascular channels within the fibrotic intima. The affected vessel is a medium sized artery within the subpericardial adipose tissue (Movat's pentachrome stain. Original magnification $\times 130$)

arteries of various caliber. The fibrous material formed thin laminar deposits that were often completely covered by endothelium that had proliferated. Fresh fibrinous deposition could be observed side by side with foci of organizing fibrosis of the intima. In some instances fibrin deposition and endothelial proliferation formed spurs that projected into the lumen as 'intimal bridges'. The internal elastic membrane was frequently normal in these vessels but their media was fibrotic.

Plexiform metamorphosis of vessels. This was the most striking change but it was present in only 50 per cent of cases. The affected arteritis appeared to have several lumina of different sizes lined by swollen endothelial cells. The wall of such vessels was almost totally replaced by fibrous tissue. Focal deposition of fibrin on the endothelial surface and endothelial polyps or projections could still be seen in vessels with advanced plexiform changes (Fig 2).

Medial sclerosis. Replacement of the medial muscle cells by fibrous tissue was seen in several arteries and arterioles in every case. The fibrous tissue varied from a loose textured basophilic tissue to a densely collagenous one.

Other changes. Several other changes also

appeared in relation to the coronary vessel such as adventitial fibrosis, perivascular edema, congestion and a slight perivascular lymphocytic accumulation. These changes were considered to be non specific rather than essentially related to the condition under study.

Comments

The presence of coronary vascular lesions in a disease such as endomyocardial fibrosis is not unexpected. Although the etiology of endomyocardial fibrosis is still obscure the various theories that have been put forward relate to processes in which vascular changes may be expected. This is true for example if we consider endomyocardial fibrosis as a form of rheumatic disease⁸ or as a muscle disease⁹ or as a hypersensitivity disease of the connective tissue of the heart.⁷

In our cases the lesions in the coronary vasculature were frequent and varied. Although mistakes readily result from trying to base a dynamic interpretation on static morphologic features we should like to link the several changes observed into a unitary sequence. Because of its frequency we consider medial degeneration to be the fundamental lesion. It is probably fol-

loved by fibrosis and by the deposition of fibrin in the intima and the organizing intimal changes. The vascular lesions do not seem to be secondary to the endomyocardial fibrosis itself since they also appeared in uninvolved myocardium and in the epicardium. They do not appear to represent a primary feature of the disease since there was no correlation between them and the degree of endomyocardium scarring. It seems to us to be probable that the changes that result in the intimal and medial fibrosis may be caused by the same still obscure factor or factors that cause the endomyocardial fibrosis. The arterial lesions could lead to further myocardial damage and thus be a potentially important element in the pathogenesis of the disease.

Medial degeneration in coronary arteries with subsequent sclerosis and calcification has been recognised in man and in experimental animals under several different conditions.⁹ Usually such medial change is followed by intimal fibrosis but repeated deposition of fibrin and the plexiform metamorphosis of the vessels do not seem to be features of the vascular changes in these cases. Medial degeneration of the smaller coronary vessels has also been described in a variety of hereditary musculo-skeletal and neuromuscular syndromes.¹⁰ As Jamieson¹¹ suggested the possibility that a hereditary medial necrosis of small coronary arteries is responsible for some obscure cardiomyopathies deserves further study.

Medial degeneration of the smaller branches of the coronary arteries is an important finding in dogs that have been maintained for long periods on a diet deficient in magnesium.¹ A similar lesion has been found in alcoholic cardiopathy.¹² It may be significant that hypomagnesaemia has been noted in cases of chronic alcoholism and of hepatic cirrhosis.¹³ Similarly, endocardial necrosis that heralds with endomyocardial scarring has been found in rats treated with corticoids and sodium salts; such necrosis was prevented by the prophylactic administration of magnesium.¹ At present unfortunately nothing seems to be known about the magnesium content of the blood of patient with endomyocardial fibrosis.

Regardless of their obscure aetiology we consider those obstructive coronary arterial lesions that were a prominent histopathological feature of our cases of endomyocardial fibrosis as a pathogenetic element in the progression of this disease.

Summary

A series of changes affecting the coronary arteries was found during a postmortem histopathological study of 12 consecutive cases of endomyocardial fibrosis in Brazil. These changes seem to represent a spectrum which includes medial degeneration, medial sclerosis, organizing fibrin deposition on the intimal surface and the formation of plexiform lesions. Medium sized and small arteries and arterioles within the fibrotic endomyocardial areas were mainly affected but changes were also seen in other areas of the myocardium and in the epicardium. Lesions were not found in the vasculature outside the heart.

The lesions in the coronary vasculature resemble morphologically the basic process of fibrosis that affects the endocardium and adjacent myocardium; they are more over an important feature of the histopathological picture of the disease and may play a role in the pathogenesis of the accompanying myocardial damage.

REFERENCES

- 1 Andrade Z A and Guimaraes A L. Endomyocardial fibrosis in Salvador Bahia, Brazil. *Br Heart J* 26: 813, 1964.
- 2 Adriaens D G. Endomyocardial fibrosis of the right ventricle. *Quart J Med* 31: 1, 1967.
- 3 Connor D H, Somers K, Hunt W S R, Wainman W C and D'Arbellet P C. Endomyocardial fibrosis in Uganda (Davies disease). *Am Heart J* 74: 687, 1967 and 75: 107, 1968.
- 4 Davies J N P and Bill J D. The pathology of endomyocardial fibrosis in Uganda. *Br Heart J* 17: 337, 1955.
- 5 Pirry E H O and Abrahams D G. The natural history of endomyocardial fibrosis. *Quart J Med* 74: 383, 1965.
- 6 Edington G M and Jackson J G. The pathology of heart muscle disease and endomyocardial fibrosis in Nigeria. *J Pathol* 86: 333, 1963.
- 7 McKinney M. A comparative histological study of endomyocardial fibrosis and cardiomyopathy of unknown origin. *Pathol Microbiol* 23: 70, 1970.
- 8 Shipner A G. Endomyocardial fibrosis and rheumatic heart disease. *Lancet* 1939: 1966.
- 9 Friesen R W and Fidler R S. Concomitant

- medial sclerosis of the coronary artery. *Am Heart J* 7:133, 1931
- 10 James T N. Pathology of small coronary arteries. *Am J Cardiol* 20:679, 1967
- 11 James T N. An etiologic concept concerning the obscure myocardopathies. *Proc Cardio-vasc Dis* 7:43, 1964
- 12 Wener J, Pintar H, Simon M A, Motola R, Friedman R, Myrman A and Schueer K. The effect of prolonged hypomagnemicemia on the cardiovascular system in young dogs. *Am Heart J* 67:221, 1964
- 13 Pintar H, Wolanski B M and Gubbas I R. Alcoholic cardiomyopathy. *Can. Med Assoc J* 93:103, 1965
- 14 Martin H E and Bruer F K. Magnesium studies in the cirrhotics and alcoholics. *Proc R Soc Med* 55:917, 1961
- 15 Seyle H. Experimental production of endomyocardial fibrosis. *Lancet* 1:1331, 1958

Comparison of pulmonary wedge and left atrial pressure in man

Abe Walston II MD

N Eugene Kendall MD

Durham N C

As early as 1906 it was observed that pressure change in the left atrium and pulmonary venous system could be transmitted retrograde to the end arteries of the lungs. Pulmonary wedge pressures were first described by Hellum and associates¹ in 1948. The development of left heart catheterization allowed direct measurement of left atrial pressure and provided a means for direct correlation of pulmonary wedge and left atrial pressures. On the basis of these studies pulmonary wedge pressures have been used as synonymous with the left atrial pressure in the calculation of mitral valve areas and other hemodynamic indexes. The hemodynamic evaluation of patients with acute myocardial infarction has led to the use of pulmonary wedge pressure as an index of left ventricular filling pressure.^{2,3} This recent use of pulmonary wedge pressure has placed new emphasis on defining the precise correlation of mean pulmonary wedge and mean left atrial pressure. Our review of the literature fails to reveal a comparison of pulmonary wedge and left atrial pressures in a large number of patients. This study was designed to compare retrospectively mean pulmonary wedge and

mean left atrial pressure in a large number of patients with and without heart disease.

Methods

The records of 700 patients obtained at cardiac catheterization were analyzed. The hemodynamic data were obtained as part of the diagnostic cardiac catheterization at Duke University Medical Center and Durham Veterans Administration Hospital between 1959 and 1972. The patient groups included normals (23), coronary artery disease (54), mitral stenosis (159), mitral insufficiency (50), aortic stenosis (65), aortic insufficiency (39), and patients with more than one valvular lesion (310). Each patient underwent right and left heart catheterization with left atrial pressure measured directly by a transeptal left atrial puncture. Pulmonary wedge pressure was repeated multiple times to insure that correct values were obtained. Withdrawal of 95 per cent saturated hemoglobin and snapping of the catheter tip on withdrawal of the wedged catheter was used as evidence of a correct wedge position. Whenever possible pulmonary wedge pressures and left atrial pressures were recorded simultaneously on equally cali-

From the Department of Medicine (Dr. Walston) and the Department of Cardiology (Dr. Kendall) Duke University Medical Center and the Veterans Administration Medical Center, Durham, N.C. 27705.

This work was supported in part by United States Public Health Service Contract N01-05756 (Contract N01-05756) from the National Heart, Lung, and Blood Institute.

Received for publication Sept. 15, 1972.

Revised manuscript received Dec. 11, 1972. H. Room C-200, Veterans Administration Hospital, Durham, N.C. 27705.

Table I Comparison of mean values, ranges, and standard deviations (mm Hg) for mean pulmonary wedge and mean left atrial pressures for each of the seven patient groups and for all cases

| Group | N | Mean (mm Hg) | Range (mm Hg) | Standard deviation (mm Hg) | Correlation coefficient* |
|--------------------------|-----|-----------------|------------------|----------------------------------|-----------------------------|
| 1 Normals | 23 | | | | |
| PW† | | 7.0 | 2-15 | ±0.6 | |
| LA | | 6.4 | 2-12 | ±0.4 | 0.93 |
| 2 Coronary disease | 54 | | | | |
| PW | | 9.6 | 3-31 | ±0.5 | |
| LA | | 9.1 | 3-25 | ±0.5 | 0.91 |
| 3 Mitral stenosis | 159 | | | | |
| PW | | 20.4 | 7-43 | ±0.6 | |
| LA | | 19.2 | 6-48 | ±0.5 | 0.90 |
| 4 Mitral insufficiency | 50 | | | | |
| PW | | 13.7 | 4-37 | ±0.9 | |
| LA | | 12.8 | 3-38 | ±1.0 | 0.93 |
| 5 Aortic stenosis | 65 | | | | |
| PW | | 11.0 | 4-41 | ±0.8 | |
| LA | | 10.8 | 2-42 | ±0.9 | 0.91 |
| 6 Aortic insufficiency | 39 | | | | |
| PW | | 13.5 | 4-36 | ±1.1 | |
| LA | | 12.9 | 3-28 | ±1.0 | 0.90 |
| 7 Mixed valvular disease | 310 | | | | |
| PW | | 16.0 | 3-43 | ±0.4 | |
| LA | | 15.9 | 2-42 | ±0.4 | 0.93 |
| 8 All cases | 700 | | | | |
| PW | | 15.4 | 2-43 | ±0.3 | |
| LA | | 15.0 | 2-48 | ±0.3 | 0.93 |

*Correlation coefficients are also given for the comparison of mean pulmonary wedge and mean left atrial pressures.

†Abbreviations: PW = pulmonary wedge; LA = left atrial; N = number of patients.

brated gauges. Pressures were measured with Statham P23Db strain gauges and recorded on an Electronics for Medicine DR8 recorder. Mean pressures were obtained by electronic integration of the phasic pressures. Pressures used in this study were measured when the heart rate was below 100 beats per minute. No additional studies were carried out to determine the possible effect of heart rate on the relation between wedge and left atrial pressure. Standard linear regression techniques were used for evaluation of the data.⁴

Results

Table I shows the mean, range, standard deviation, and correlation coefficient for the mean pulmonary wedge and mean left atrial pressure for each of the patient groups. Despite a wide variation in the

range of wedge and left atrial pressures, the correlation coefficients for each of the patient groups were found to be nearly identical. Fig. 1 shows a plot of mean pulmonary wedge versus mean left atrial pressure for all patients. Linear regression analysis reveals a correlation coefficient of $r = 0.93$ with a standard error of ± 3.0 mm Hg. Note that the degree of scatter increases as the wedge and left atrial pressures rise.

Table II and Fig. 2 reveal a comparison of mean pulmonary wedge and mean left atrial pressure for all of the patients. For this comparison the data were grouped according to a 5 mm Hg increment in mean wedge pressure. The mean difference for each of these groups was computed as the difference between mean left atrial pressure and mean pulmonary wedge pressure. At wedge pressures below 25 mm Hg there

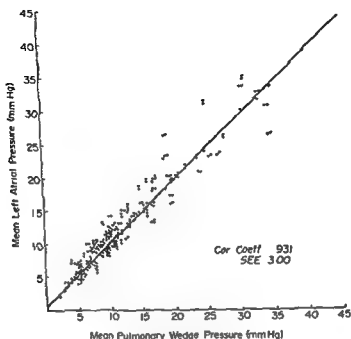


Fig. 1 Plot of mean left atrial pressure versus mean pulmonary wedge pressure (mm Hg) for all patients. The regression line was $y = 0.93x + 0.57$ and the correlation coefficient was $r = 0.93$.

were no significant differences between mean wedge and mean left atrial pressure. In wedge pressure groups 26 to 30, 31 to 35 and >35 mm Hg mean left atrial pressure was significantly different from mean wedge pressure ($P < 0.05$). The 95 per cent confidence limits of the prediction of mean left atrial pressure from mean wedge pressure for each of these groups are shown in Fig. 2. Note that with a mean wedge pressure of 10 mm Hg or less the error in predicting left atrial pressure is only ± 2 mm Hg. However the error increases considerably when the mean wedge pressure is higher.

Discussion

Hellems and colleagues¹ reported experimental evidence that the pressure recorded by a catheter wedged in an end pulmonary artery closely approximated pulmonary capillary pressure. Werko and associates², Connolly and co-workers³, Fjeps and colleagues⁴ and others^{5,6} found good correlation between the level of pulmonary wedge and left atrial pressure. Other investigators⁷⁻¹¹ in human and animal stud-

ies found poor correlation between wedge and left atrial pressure. However these early comparisons were performed with left atrial pressures measured in a variety of ways including measurement by way of atrial septal defect¹² by left atrial puncture during open heart surgery,¹³ measurement by transbronchial approach¹⁴ and measurement by transthoracic approach.^{15,16} Werko and co-workers² demonstrated that balloon occlusion of the pulmonary artery proximal to the wedged catheter does not alter the pressure recorded at the tip of the catheter demonstrating the left-sided origin of the pulmonary wedge pressure. Ankeney¹⁷ showed that phasic variations of left atrial pressure waves were transmitted retrograde to the wedged catheter but only mean left atrial pressure was accurately reflected in the wedge pressure tracings. Hemodynamic studies on patients with acute myocardial infarction have directed new emphasis on the ability of pulmonary wedge pressure to predict left atrial pressure which is used as an index of left ventricular filling pressure.¹⁸ Pulmonary wedge pressures (using

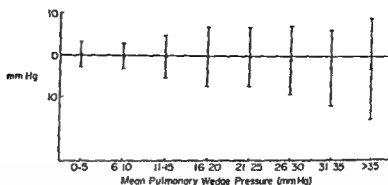


Fig 2 Plot of mean difference in mm Hg of left atrial pressure from mean wedge pressure (mm Hg) in groups of 5 mm Hg of mean wedge pressure. Points above and below the zero line signify mean left atrial pressure greater and less than respectively mean pulmonary wedge pressure. Brackets surrounding these mean points signify the 95 per cent confidence limits. (The 95 per cent confidence limits of the data were determined by multiplying the standard error of the mean by the appropriate T value).⁴

Table II. Mean pulmonary wedge pressure vs mean left atrial pressure

| Pulmonary wedge pressure group (mm Hg) | N* | Mean difference (mm Hg) | Standard error of the mean (mm Hg) |
|--|-----|-------------------------|------------------------------------|
| 0-5 | 39 | 0 | ±0 |
| 6-10 | 207 | 0 | ±0 |
| 11-15 | 163 | 0 | ±0 |
| 16-20 | 111 | 0 | ±0 |
| 21-25 | 84 | 0 | ±0 |
| 26-30 | 52 | -1† | ±1 |
| 31-35 | 28 | -3† | ±1 |
| >35 | 16 | -3† | ±1 |

*N = number of patients in each 5 mm Hg incremental group.

†Signifies that the mean difference was statistically significant ($P < 0.05$).

the Swan Ganz catheter) can be monitored at the bedside and if valid can provide vital data on left ventricular filling pressure.^{20, 21} Since left ventricular filling pressure is frequently elevated in acute myocardial infarction, correlation of pulmonary wedge with left atrial pressure must be done over a wide range of pressures.

The plot of mean wedge versus mean left atrial pressure (Fig 1) demonstrates little scatter around the regression line until the pressures exceed 10 mm Hg. The near identity of the mean differences and the small standard errors between wedge and left atrial pressure suggest that mean pulmonary wedge pressure is a moderately accurate predictor of mean left atrial pressure at these low normal wedge pressures. The predictive ability of mean wedge pressure

(described by the 95 per cent confidence limits illustrated in Fig 2) is directly related to the level of mean wedge pressure. When the wedge pressure is greater than 10 mm Hg the prediction of left atrial pressure can be shown to contain considerable error. Ventricular filling pressure and other assessments of myocardial performance and valve areas cannot validly be calculated from the wedge pressure if it exceeds 10 mm Hg. The increasing loss of correlation between wedge and mean left atrial pressure at higher pressures may in part be explained by the findings of Caro and associates²² who demonstrated that high pressures across the pulmonary capillary bed cause a change in the ratio of pulmonary arterial and pulmonary venous compliance promoting asymmetrical trans-

mission of pressure waves across the pulmonary vascular bed

The present study was done in a retrospective manner and has all the errors in measurement which are inherent in the day-to-day operation of a diagnostic cardiac catheterization laboratory. Thus it might be argued that meticulous attention to detail in pressure recording might yield a closer correlation. However it is with this level of expertise in pressure measurement that hemodynamic data is collected on a routine basis.

Summary

Values of mean pulmonary wedge and mean left atrial pressure were compared in 700 patients including normals and patients with a variety of heart diseases. An overall correlation coefficient $r = 0.93$ was found between mean wedge and mean left atrial pressure and this correlation was unrelated to the etiology of the heart disease. No significant difference between the average value of wedge and left atrial pressure was found until the wedge pressure exceeded 25 mm Hg. In patients with mean wedge pressures of 10 mm Hg or less left atrial pressure could be predicted with an accuracy of ± 2 mm Hg (95 per cent confidence limits). However at higher wedge pressures the prediction of mean left atrial pressure was subject to considerable error. Thus caution should be employed in using mean wedge pressure as an accurate index of mean left atrial pressure unless the wedge pressure is within the low normal range.

The authors gratefully acknowledge the technical assistance of Judith C. Pembert, Ph.D., The Department of Medical Illustrations of the Durham Veterans Administration Hospital also rendered valuable support in the preparation of this paper.

The secretarial assistance of Mrs. Rosa M. Ethridge and Mrs. Brenda Haley is hereby acknowledged. Statistical analysis was performed by Philip McHale, Ph.D. The authors are also grateful to Dr. Yuhong Hong of the Duke Cardiovascular Laboratory for making the catheterization data available.

REFERENCES

1. Hellem H. K., Haynes F. W., Dexter L. and Kinney T. D. Pulmonary capillary pressure in animals estimated by venous and arterial catheterization. *Am J Physiol* 150: 98, 1948.
2. Forrester James S., Diamond George

- McHugh Thomas J. and Swan H. J. C. Filling pressures in the right and left sides of the heart in acute myocardial infarction. *N Engl J Med* 283: 190, 1971.
3. Rackley C. E. and Russell R. O. Left ventricular function in acute myocardial infarction and its clinical significance. *Circulation* 45: 231, 1972.
4. Snedecor G. W. Statistical methods. 5th ed. Ames, 1962. Iowa State University Press.
5. Hellem H. K., Haynes F. W. and Dexter L. Pulmonary capillary pressure in man. *J Appl Physiol* 2: 21, 1949.
6. Werko L., Larnaukas E., Eliasson H., Lagerlof H., Senning A. and Thomasson B. Further evidence that the pulmonary capillary venous pressure pulse in man reflects cyclic pressure changes in the left atrium. *Circ Res* 1: 337, 1953.
7. Connolly D. C., Tompkins R. G., Lev R., Kirklin J. W. and Wood E. H. Pulmonary artery wedge pressures in mitral valve disease: relationship to left atrial pressures. *Mayo Clin Proc* 28: 72, 1953.
8. Connolly D. C., Kirklin J. W. and Wood E. H. The relationship between pulmonary artery wedge pressure and left atrial pressure in man. *Circ Res* 2: 434, 1954.
9. Epps R. G. and Adler R. H. Left atrial and pulmonary capillary venous pressures in mitral stenosis. *Br Heart J* 15: 798, 1953.
10. Wilson R. H., McHenna W. T., Johnson F. E., Jensen N. K., Mazzitello W. F. and Dempsey M. S. The significance of the pulmonary arterial wedge pressure. *J Lab Clin Med* 42(3): 408, 1953.
11. Wilson R. H., Joseph W. and Dempsey M. The interrelations of the pulmonary arterial and venous pressures. *Circ Res* 3: 3, 1955.
12. Samet P., Litwak R. S., Bernstein W. H., Fiere E. M. and Silverman L. M. Clinical and physiologic relationships in mitral valve disease. *Circulation* 19: 517, 1959.
13. Haddy F. J., Alden J. F., Ferrin A. L., Hannon D. W., Adams W. L. and Baronovsky I. D. An evaluation of wedge pressures in dogs under conditions of normal and elevated pulmonary vascular pressures. *Circ Res* 1: 137, 1953.
14. Murphy J. P. Inaccuracy of wedge pressure as an index of pulmonary capillary pressure. *Circulation* 27: 199, 1958.
15. Bernstein W. H., Fiere E. M., Laslo M. H., Samet P. and Litwak R. S. The interpretation of pulmonary artery wedge (pulmonary capillary) pressures. *Br Heart J* 22: 37, 1960.
16. Luchsinger P. C., Seipp H. W. and Patel D. J. Relationship of pulmonary artery wedge pressure to left atrial pressure in man. *Circ Res* 11: 315, 1967.
17. Calazel P., Gerard H., Daley R., Draper A., Foster J. and Bing R. J. Physiological studies in congenital heart disease. VI. A comparison of the right and left auricular capillary and pulmonary artery pressures in nine patients.

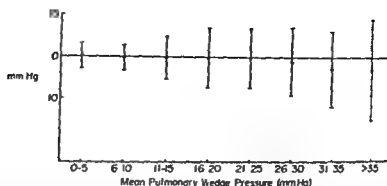


Fig. 2 Plot of mean difference in mm Hg of left atrial pressure from mean wedge pressure (mm Hg) in groups of 5 mm Hg of mean wedge pressure. Points above and below the zero line signify mean left atrial pressure greater and less than respectively mean pulmonary wedge pressure. Brackets surrounding the mean point signify the 95 per cent confidence limits. (The 95 per cent confidence limits of the data were determined by multiplying the standard error of the mean by the appropriate T value).⁴

Table II Mean pulmonary wedge pressure vs. mean left atrial pressure

| Pulmonary wedge pressure group (mm Hg) | N* | Mean difference (mm Hg) | Standard error of the mean (mm Hg) |
|--|-----|-------------------------|------------------------------------|
| 0-5 | 39 | 0 | ±0 |
| 6-10 | 207 | 0 | ±0 |
| 11-15 | 163 | 0 | ±0 |
| 16-20 | 111 | 0 | ±0 |
| 21-25 | 84 | 0 | ±0 |
| 26-30 | 52 | -1† | ±1 |
| 31-35 | 28 | -3† | ±1 |
| >35 | 16 | -3† | ±1 |

*N = number of patients in each 5 mm Hg incremental group

†Signifies that the mean difference was statistically significant ($P < 0.05$)

the Swan Ganz catheter) can be monitored at the bedside and if valid can provide vital data on left ventricular filling pressure.^{20,21} Since left ventricular filling pressure is frequently elevated in acute myocardial infarction, correlation of pulmonary wedge with left atrial pressure must be done over a wide range of pressures.

The plot of mean wedge versus mean left atrial pressure (Fig. 1) demonstrates little scatter around the regression line until the pressures exceed 10 mm Hg. The near identity of the mean differences and the small standard errors between wedge and left atrial pressure suggest that mean pulmonary wedge pressure is a moderately accurate predictor of mean left atrial pressure at these low normal wedge pressures. The predictive ability of mean wedge pressure

(described by the 95 per cent confidence limits illustrated in Fig. 2) is directly related to the level of mean wedge pressure. When the wedge pressure is greater than 10 mm Hg the prediction of left atrial pressure can be shown to contain considerable error. Ventricular filling pressure and other assessments of myocardial performance and valve areas cannot validly be calculated from the wedge pressure if it exceeds 10 mm Hg. The increasing loss of correlation between wedge and mean left atrial pressure at higher pressures may in part be explained by the findings of Caro and associates² who demonstrated that high pressures across the pulmonary capillary bed cause a change in the ratio of pulmonary arterial and pulmonary venous compliance promoting asymmetrical trans-

The progression of coronary atherosclerotic disease as assessed by cine-coronary arteriography

Robert R. Henderson M.D.
George G. Rowe M.D.
Madison, Wis.

The development of cine-coronary arteriography has enabled the physician to define obstruction of the coronary arterial system and to demonstrate the rate of progression of atherosclerotic coronary artery disease in individual patients. The present study was undertaken to evaluate arteriographic progression of coronary disease in 64 patients, 7 of whom were treated with partial ileal bypass surgery and to correlate progression with clinical and biochemical parameters.

Groups studied and methods of study

Two groups of subjects were studied. The first group consisted of 8 patients, 7 men and one woman with hyperlipidemia and severe coronary atherosclerosis documented by coronary arteriography who underwent partial ileal bypass surgery. One subject died early, leaving 6 men and one woman for restudy. Postoperative arteriograms were performed in all patients (the mean postoperative interval was 25 months) in order to follow their coronary arterial disease. The second group of 57 patients, 54 men and 3 women underwent repeat coronary arteriography after an

interval of one to 54 months (mean 14 months) to evaluate revascularization procedures or because of progressive symptoms. These two groups are considered separately throughout the study.

Age, height, weight, history of previous myocardial infarction (substantiated by electrocardiographic and/or serum enzyme changes), family history of coronary artery disease and cigarette consumption were noted. In 63 subjects the presence or absence of glucose intolerance was established by a glucose tolerance or a two hour post prandial blood sugar test following a standard 100 Gm oral glucose load. Fasting serum cholesterol levels in all patients and triglyceride levels in 36 patients were determined by standard laboratory methods. Lipoprotein phenotyping was done in 50 patients by the method of Fredrickson and associates.¹ Chest roentgenograms were reviewed to establish the presence or absence of cardiomegaly as determined by the cardiothoracic ratio (greatest transverse diameter of the heart greater than one half the greatest transverse diameter of the chest). A history of myocardial infarction in the interval between coronary

From the Cardiovascular Research Laboratory, Department of Medicine, University of Wisconsin Medical School, Madison, Wis.

This research was supported in part by the U.S. Health Statistics, Public Health Service, National Institutes of Health Grants HL-07754 and HL-5364.

Received for publication Sept. 25, 1972.

Reprint requests to George G. Rowe, M.D., Cardiovascular Research Laboratory, 400 North Lake Street, Room 523, Madison, Wis. 53706.

- with auricular septal defect *Johns Hopkins Med J* 88 20 1951
- 18 Werko L Varnauskas E Eliash H and Thomasson B The influence of the pulmonary arterial pressure on the pulmonary capillary venous pressure in man *Circ Res* 1:340 1953
- 19 Ankenney J I Further experimental evidence that pulmonary capillary pressures do not reflect cyclic changes in left atrial pressure (mitral lesions and pulmonary embolism) *Circ Res* 1:58 1953
- 20 Ganz W W Forrester J S Chonette, D Donoso R and Swan H J C A new flow directed catheter technique for measurement of pulmonary artery and capillary wedge pressure without fluoroscopy *Am J Cardiol* 26:66 1970
- 21 Cohn J N Khatri I M and Hamosh P Bedside catheterization of the left ventricle *Am J Cardiol* 25 66 1970
- 22 Caro C G Bergel D H and Seed W A Forward and backward transmission of pressure waves in the pulmonary vascular bed of the dog *Circ Res* 20 185 1967

Table 1 Clinical and biochemical data for 7 patients in Group I (partial ileal bypass)

| Datum | Value |
|--|---------------------------|
| Age | 35 ± 10 |
| Sex (% male) | 86% |
| Weight (lb) | 168 ± 21 |
| Height (in) | 67 ± 2 |
| Smokers | 85% |
| Family history of coronary disease | 57% |
| Carbohydrate intolerance | 0 |
| Serum cholesterol (mg %) | |
| Preoperative | 452 ± 99 |
| Postoperative | 236 ± 42 ($p < 0.001$)† |
| Serum triglyceride (mg %) (n = 5) | |
| Preoperative | 211 ± 107 |
| Postoperative | 112 ± 20 (NS) |
| Past myocardial infarction | 27% |
| Interim myocardial infarction | 27% |
| Card omegaly | 0 |
| Left ventricular end-diastolic pressure‡ | 17 ± 8 |
| Aortic mean pressure | 103 ± 14 |
| Interval between partial ileal bypass and repeat coronary arteriogram (mo) | 25 ± 7 |

M ± st. dev. d. lat. n.

†St. d. test and N.S. d. one $p > 0.05$

‡Left ventricular end-diastolic pressure

preoperatively versus 273 ± 95 mg per 100 ml postoperatively) in 5 patients with arteriographic progression ($p < 0.01$). Serum triglyceride levels measured in 5 patients decreased by 47 per cent but this change failed to attain statistical significance (Table I Fig 2). One example of clear-cut progression from this group of subjects is seen in Fig 1 A and B.

Group II (all arteries) Progression was noted in 37 patients (63 per cent). The interval between coronary arteriograms was significantly longer ($p < 0.001$) in patients with arteriographic progression (17 ± 14 months) than in patients without arteriographic progression (9 ± 12 months). There were no significant differences between the two groups in the other parameters measured (Table II). Progression was noted in 37 sites in 30 arteries. Seven of these sites were in grafted arteries proximal to the saphenocoronary anastomosis. No significant progression was seen in a grafted artery distal to the saphenocoronary anastomosis.

Group III (excluding revascularized arteries) If vessels subjected to direct revascularization procedures are excluded

from the evaluation of progression 19 patients (33 per cent) showed progression involving 40 sites and 33 arteries. The difference in the interval between coronary arteriograms in patients with arteriographic progression (23 ± 15 months) versus those without arteriographic progression (8 ± 9 months) is more striking with bypassed arteries excluded ($p < 0.001$). Patients with progressive disease also had significantly higher ($p < 0.01$) mean serum cholesterol levels (Table III). The amount of progression in per cent was divided by the number of months over which the change occurred to derive a rate of progression. This rate was then related to the serum cholesterol to determine the correlation coefficient between the two variables but the r value ($p > 0.1$) indicated no significant relation between the two variables. Serum triglyceride levels and lipoprotein phenotyping were obtained in only 5 of 19 patients with and 24 of 28 patients without arteriographic progression making statistical comparison of these parameters impractical. No significant differences were found between patients with and without arteriographic



Fig 1 Right coronary arteriograms of Patient R. Z. showing clear progression. Partial ileal bypass was performed in October 1966.

arteriograms substantiated by electrocardiographic and/or serum enzyme changes, was noted. At cardiac catheterization pressures were recorded in the left ventricle and aorta with special reference to left ventricular end diastolic and aortic mean pressure. Coronary arteriography was performed by the Sones technique² in most instances after sublingual administration of 0.4 mg. of nitroglycerin and/or 5 mg. of isosorbide dinitrate and recorded on 35 mm. film with grid pulse control x-ray equipment. Views were taken of the left coronary artery in 60, 45 and 30 degrees of left anterior oblique and 30 degrees of right anterior oblique; the right coronary artery was filmed in 45 degrees of left anterior oblique and 30 degrees of right anterior oblique. Aortocoronary saphenous vein grafts were filmed in 45 degrees of left anterior oblique and 30 degrees of right anterior oblique.

All arteriograms were independently evaluated by both of the authors. When the interpretations were not in agreement the arteriograms were reviewed jointly and a compromise interpretation reached.

Arteriographic criteria for progression were total occlusion of a previously open vessel, an increase of 25 per cent in obstruction of the vessel lumen by a pre-existing lesion, or appearance of a lesion obstructing

at least 25 per cent of the lumen of a previously normal vessel.

Forty-one patients underwent direct coronary arterial revascularization procedures to 57 vessels in the interval between coronary arteriograms. Thirty-three patients had 49 saphenous vein grafts, 5 of which were combined with an endarterectomy. One patient had an internal mammary aortocoronary graft. Two patients underwent endarterectomy alone and 3 had an endarterectomy with a patch graft. Twelve patients underwent Vinberg procedures, 3 in conjunction with direct coronary artery surgery.

Statistical analysis was performed using the Student *t* test, the chi square test, or the simple correlation coefficient when appropriate.

Results

Group I (ileal bypass). Five of 7 patients (71 per cent) showed arteriographic progression; one did not have arteriographic progression and in the other the arteriograms were not technically adequate for evaluation (Fig 1). Serum cholesterol levels decreased by 48 per cent (452 ± 99 mg per 100 ml preoperatively versus 236 ± 42 mg per 100 ml postoperatively) in the group as a whole ($p < 0.001$) and 41 per cent (466 ± 117 mg per 100 ml

progression in the other parameters measured (see Table III)

Location of lesions The location of obstructive lesions showed a marked predilection for the proximal third of the coronary arteries. In Group II as a whole at the time of the initial arteriogram lesions were in proximal locations in 91 per cent for the left anterior descending, 81 per cent for the circumflex and 65 per cent for the right coronary artery (Table IV). New or progressive lesions showed this proximal predilection even when bypassed arteries were excluded. In the left anterior descending artery 92 per cent of new or progressive lesions involved the proximal third of the vessel. This was not as striking in the circumflex artery in which 56 per cent of the new or progressive lesions involved the proximal third, or the right coronary artery in which 47 per cent of the lesions were in the proximal third of the vessel (Table V). Only two patients had arteriographic progression of the left main coronary artery. One developed 50 per cent obstruction in the vessel which was only slightly involved previously and in the other a 90 per cent obstruction of the left main coronary artery progressed to total occlusion.

In an effort to determine whether proximal location or high grade luminal obstruction predisposes to progression the distribution and severity of the lesions at the time of initial arteriography in those whose lesions progressed was compared with those whose lesions remained stable. As seen in Table IV the distribution of lesions throughout the vessels was not different for the two groups. The mean degree of luminal obstruction by proximally located lesions at the time of initial coronary arteriography was not significantly different in patients with and without arteriographic progression (Table VI).

Surgical results Forty three of the 49 saphenous vein bypass grafts and the single internal mammary bypass graft were patent at restudy for a patency rate of 86 per cent. All arteries upon which endarterectomy alone or endarterectomy and patch graft was performed were patent.

Of the 50 arteries upon which bypass surgery was performed 13 were completely occluded preoperatively. All bypass grafts to these 13 arteries were patent at the

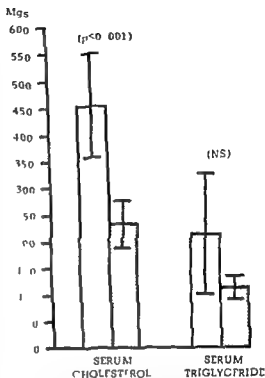


Fig 2 Preoperative (open bars) and postoperative (stippled bars) serum cholesterol and triglyceride levels in Group I patients (partial distal bypass)

Postoperative study Of the remaining 37 arteries 22 appeared to be completely occluded proximal to the saphenocoronary anastomosis postoperatively. Patients with and without proximal arterial occlusion did not differ significantly in age, serum cholesterol, per cent of preoperative occlusion (> 80 per cent both groups) or interval between surgery and postoperative arteriogram. Four patients had the unfortunate combination of a nonpatent bypass graft and complete occlusion of the grafted artery.

Only 6 of the 34 patients undergoing coronary bypass surgery had evidence of progression in nongrafted vessels. The graft patency rate in this group was 67 per cent (6 of 9) as compared to 90 per cent (37 of 41) in the patients without progression in nongrafted vessels ($p < 0.01$). The rate of artery occlusion proximal to the saphenocoronary anastomosis was not significantly different: 50 per cent (3 of 6) versus 61 per cent (19 of 31) respectively.

Table II Clinical and biochemical data for patients in Group II (bypassed arteries included)

| <i>Datum</i> | <i>Arteriographic progression (19)</i> | <i>No arteriographic progression (38)</i> |
|--|--|---|
| Age | 48 ± 9* | 49 ± 8 |
| Sex (% male) | 94% | 95% |
| Weight (lb) | 172 ± 29 | 167 ± 21 |
| Height (in) | 67 ± 3 | 68 ± 3 |
| Smokers | 78% | 81% |
| Family history of coronary disease | 56% | 52% |
| Carbohydrate intolerance | 0 | 5% |
| Serum cholesterol (mg %) | 252 ± 48 | 244 ± 43 |
| Serum triglyceride (mg %) | 170 ± 110 (17) | 160 ± 98 (14) |
| Lipoprotein phenotype | (15) | (14) |
| II | 0 | 0 |
| III | 0 | 7% |
| IV | 40% | 36% |
| Normal | 60% | 57% |
| Past myocardial infarction | 42% | 33% |
| Interim myocardial infarction | 3% | 5% |
| Cardiomegaly | 17% | 14% |
| Left ventricular end diastolic pressure (mm Hg)† | 17 ± 9 | 19 ± 8 |
| Aortic mean pressure (mm Hg) | 106 ± 22 | 100 ± 15 |
| Interval between coronary arteriograms (mo) | 17 ± 14 | 9 ± 12 |

Number of patients in parentheses

*Mean ± standard deviation

†Left ventricular end diastolic pressure

Table III Clinical and biochemical data for patients in Group II (bypassed arteries excluded)

| <i>Datum</i> | <i>Arteriographic progression (19)</i> | <i>No arteriographic progression (38)</i> | <i>p value*</i> |
|--|--|---|-----------------|
| Age | 49 ± 9† | 49 ± 8 | NS |
| Sex (% male) | 95 | 95% | NS |
| Weight (lb) | 170 ± 17 | 170 ± 28 | NS |
| Height (in) | 69 ± 3 | 68 ± 2 | NS |
| Smokers | 82% (17) | 86% (36) | NS |
| Family history of coronary disease | 63% | 54% (37) | NS |
| Carbohydrate intolerance | 0 | 5% (37) | NS |
| Serum cholesterol (mg %) | 276 ± 53 | 235 ± 47 | < 0.01 |
| Serum triglyceride (mg %) | 168 ± 103 (5) | 195 ± 102 (26) | |
| Lipoprotein phenotype | (5) | (24) | |
| II | 0 | 0 | |
| III | 0 | 4% | |
| IV | 20% | 42% | |
| Normal | 80% | 54% | |
| Past myocardial infarction | 53% | 32% | NS |
| Interim myocardial infarction | 5% | 3% | NS |
| Cardiomegaly | 16% | 16% | NS |
| Left ventricular end diastolic pressure (mm Hg)† | 18 ± 10 | 18 ± 7 | NS |
| Aortic mean pressure (mm Hg) | 105 ± 18 | 101 ± 16 | NS |
| Interval between coronary arteriograms (mo) | 23 ± 15 | 9 ± 9 | < 0.001 |

Number of patients in parentheses

*Student t test or chi square used NS denotes $p > 0.05$

†Mean ± standard deviation

‡Left ventricular end diastolic pressure

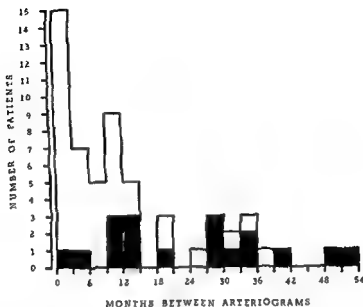


Fig 3 Interval between arteriograms in months and number of patients studied at each interval. Shaded squares represent patients with arteriographic progression.

Table VI Mean per cent of luminal obstruction by lesions in the proximal third of coronary arteries in Group II patients at the initial arteriogram. The group is divided into those patients who showed progression at repeat arteriography and those who did not.

| Vessel | Arteriographic progression (36) | No arteriographic progression (21) | p Value |
|--------------------------|---------------------------------|------------------------------------|---------|
| Left anterior descending | 75 ± 21 | 80 ± 20 | NS |
| Circumflex | 67 ± 20 | 73 ± 23 | NS |
| Right | 71 ± 22 | 71 ± 25 | NS |
| All vessels | 72 ± 21 | 77 ± 21 | NS |

NS = not significant
Mean ± standard deviation

were felt severe enough to require surgical revascularization whereas this had not been the case after the first arteriogram. Another patient developed a 50 per cent occlusion of the left anterior descending in 3 months time. The significantly higher serum cholesterol levels in patients with arteriographic progression is expected and is in agreement with Bemis and associates⁴ who studied 50 patients with repeat cine coronary arteriography (mean interval 23 months) using similar criteria for progression. The Bemis group also found elevated serum triglyceride levels to be correlated with progression. Ten of 29 patients in our

series without arteriographic progression in whom lipoprotein phenotyping was performed were found to be Type IV. Had the interval between coronary arteriograms in the patients been longer, progression may have been found.

It would seem logical that atherosclerotic obstructive coronary arterial disease would progress with time and this is supported by the significantly longer interval between arteriograms in our patients with arteriographic progression than in our patients without arteriographic progression. Thus there was progression in 9 of 13 subjects restudied after 2 years and in

Table IV Percentage of lesions involving the proximal third of the coronary arteries at the initial arteriography in Group II patients. The group is further divided into patients who showed progression at repeat arteriography and those who did not

| Artery | All patients (57) | Arteriographic progression (19) | p value* | No arteriographic progression (38) |
|--------------------------|-------------------|---------------------------------|----------|------------------------------------|
| Left anterior descending | 91 | 85 | NS | 93 |
| Circumflex | 81 | 82 | NS | 80 |
| Right | 65 | 65 | NS | 66 |

Number of patients in parentheses

*Chi square used. N. S. denotes $p > 0.05$

Table V Percentage of new or progressive lesions involving the proximal third of coronary arteries in Group II patients at repeat arteriography

| Artery | % |
|--------------------------|----|
| Left anterior descending | 92 |
| Circumflex | 56 |
| Right | 47 |

Discussion

It is not feasible to compare Groups I and II statistically because of the discrepancies in size and constituency. However, it is of interest to note that of 6 ideal bypass patients in whom coronary arteriograms were of sufficient quality for comparison, there was clear arteriographic progression in 5 patients and in 3 myocardial revascularization was done to relieve progressive angina pectoris despite a very significant reduction in serum cholesterol levels. This is in contrast with the experience of Knight and associates² who have obtained repeat coronary arteriograms in 18 patients with ideal bypass surgery and found progression in 4 patients. Clearly, information is needed on the rate of progression of coronary atherosclerosis and whether it is altered by ideal bypass.

Myocardial revascularization, especially the aortocoronary saphenous bypass procedure, has played a prominent role in the therapy of obstructive coronary arterial disease at this institution. Consequently, few patients have had repeat cinecoronary

arteriography without surgical intervention in the interim. All arteries including those subjected to surgical revascularization procedures, were initially included in the evaluation of progression (Group II). Bypassed arteries whose proximal portions were not filled by contrast material at arteriography were assumed to be occluded. However, because of flow alterations which may occur in arteries subjected to revascularization surgery, it seems reasonable to exclude these arteries from the assessment of progression (Group II, excluding revascularized arteries). Occlusion of a bypassed artery in an area of high grade stenosis proximal to the saphenocoronary anastomosis cannot be assumed to be equivalent to atherosclerotic progression in vessels not subjected to bypass surgery, especially when progression was not seen distal to the saphenocoronary anastomosis. In addition, technical features of the surgical procedure with constriction or clotting at or near the anastomotic site may lead to occlusion. Furthermore, one must consider the possibility that failure to opacify an area previously shown to be tightly stenotic in the proximal portion of a recently bypassed vessel does not prove that the area is occluded. It is difficult to explain why contrast material would flow through a tightly stenotic area when the artery proximal and distal to it are both subjected to equal pressures from the aorta, with the pressure to the distal segment transmitted via a saphenous vein graft.

In Group II, excluding revascularized arteries, clear progression was found in 19 of 57 patients (33 per cent). In one patient the lesions found at the second arteriogram

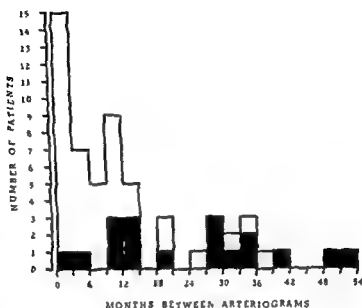


Fig. 3 Interval between arteriograms in months and number of patients studied at each interval. Shaded squares represent patients with arteriographic progression.

Table VI Mean per cent of luminal obstruction by lesions in the proximal third of coronary arteries in Group II patients at the initial arteriogram. The group is divided into those patients who showed progression at repeat arteriography and those who did not.

| Vessel | Arteriographic progression (36) | No arteriographic progression (21) | P value |
|--------------------------|---------------------------------|------------------------------------|---------|
| Left anterior descending | 75 ± 21 | 80 ± 20 | NS |
| Circumflex | 67 ± 20 | 73 ± 23 | NS |
| Right | 71 ± 22 | 71 ± 25 | NS |
| All vessels | 72 ± 21 | 77 ± 21 | NS |

NS, number of patients in parentheses.
Mean ± standard deviation.

were felt severe enough to require surgical revascularization whereas this had not been the case after the first arteriogram. Another patient developed a 50 per cent occlusion of the left anterior descending in 3 months time. The significantly higher serum cholesterol levels in patients with arteriographic progression is expected and is in agreement with Bemis and associates⁴ who studied 50 patients with repeat cine coronary arteriography (mean interval 23 months) using similar criteria for progression. The Bemis group also found elevated serum triglyceride levels to be correlated with progression. Ten of 29 patients in our

series without arteriographic progression in whom lipoprotein phenotyping was performed were found to be Type IV. Had the interval between coronary arteriograms in the patients been longer, progression may have been found.

It would seem logical that atherosclerotic obstructive coronary arterial disease would progress with time and this is supported by the significantly longer interval between arteriograms in our patients with arteriographic progression than in our patients without arteriographic progression. Thus there was progression in 9 of 13 subjects restudied after 2 years and in

only 6 of 36 restudied at one year or less (Fig. 3). We have no explanation for the failure of Bemis and co workers to find progression related to the time elapsed between arteriograms.

The predilection of atherosclerotic obstructive coronary artery disease for proximal portions of the coronary arterial tree in these patients is in agreement with post mortem findings.^{1,2} As expected, new and progressive lesions as demonstrated here by arteriography shared this proximal predilection. Proximal location and mean degree of luminal obstruction by lesions at initial arteriography were not related to progression.

The 59 per cent rate of occlusion of coronary arteries proximal to the saphenocoronary anastomosis is disturbing especially when combined with a nonpatent bypass graft. Because of the 86 per cent graft patency rate, this was a relatively uncommon finding in this series. Our data do not suggest any parameters helpful in predicting which arteries will undergo proximal occlusion. They do suggest, however, that patients with progressive disease in vessels not grafted will have lower saphenous vein graft patency rates than those without progressive disease in non-grafted vessels.

Conclusion

Seven of 8 survivors with hyperlipidemia and obstructive coronary arterial disease who underwent ileal bypass surgery had repeat coronary arteriograms. Clear progression of coronary disease was found in 5 patients, despite a significant reduction in serum cholesterol levels. Fifty-seven patients were studied by repeat coronary arteriography to evaluate progressive symptoms or revascularization procedures. Progression was seen in 19 patients (33 per cent). The interval between arteriograms was significantly longer and the serum cholesterol levels significantly higher in patients with progression.

Fifty bypass grafts were performed in 34 patients with a patency rate of 86 per cent. Apparent occlusion of the coronary artery proximal to the saphenocoronary anastomosis was seen in 59 per cent of grafted arteries and the reliability of this observation is discussed.

REFERENCES

1. Fredrickson D S, Levy R J and Lees R E. Fat transport in lipoproteins—An integrated approach to mechanisms and disorders. *N Engl J Med* 276:32-94, 148-273, 1967.
2. Sones F M Jr. Cine coronary arteriography. *Mod Concepts Cardiovasc Dis* 31:735, 1967.
3. Knight L, Scheibel R, Amplatz H, Varco R L and Buchwald H. Partial ileal bypass: A radiographic appraisal of the Minnesota Study. *Circulation* 44(Suppl. II):705, 1971.
4. Bemis C E, Eber L M, Kemp H G and Gorlin R. Cine arteriographic progression of coronary disease: Clinical and metabolic correlates. *Circulation* 44(Suppl. II):147, 1971.
5. Saphur O, Priest W S, Hamburger W W and Katz L N. Coronary arterio sclerosis, coronary thrombosis and the resulting myocardial changes. *AM HEART J* 10:567-767, 1935.
6. Mönckeberg J G, in Henke F and Lubarsch O, editors. *Handbuch der speziellen pathologischen Anatomie und Histologie: Herz und Gefäße*. II. Berlin 1924. Springer Verlag.
7. Kirch E. *Pathologie des Herzens*. Ergebn. Allg. Path. 22:1, 1927.
8. Levine H D and Brown C L. Coronary thrombosis: Its various clinical features. *Medicine* 8:245, 1929.
9. Winternitz M D, Thomas R M and LeCompte P M. *The biology of arteriosclerosis*. Springfield, Ill. 1938. Charles C Thomas Publisher.
10. Fulton W F M. *The coronary arteries*. Springfield, Ill. 1965. Charles C Thomas Publisher.
11. Podbard S. Physical factors in the progression of stenotic vascular lesions. *Circulation* 17:410, 1958.
12. Schlesinger M J and Zoll P M. Incidence and localization of coronary artery occlusions. *Arch Path* 30:178, 1949.
13. Biroldi G and Scornazzoni G. *Coronary circulation in the normal and the pathologic heart*. Washington, 1967. Office of the Surgeon General, Department of the Navy.

Bifascicular block: A clinical and electrophysiologic study

Dorothy Lunstadt MD

Manohar Punja MD

Norman Cagin MD*

Pa-Fernandez MD

Barrie Levitt MD

Yusuf Z Yuceoglu MD FACC

New York NY

In the last ten years the electrocardiographic features of bifascicular block have become well defined and their pathological correlations have been described.¹⁻⁵

The clinical implications of this abnormality are also better understood. Thus it is known that bifascicular block precedes the development of complete heart block in about 50 per cent of patients with complete heart block.⁶ Since in patients with bifascicular block atrioventricular conduction depends on the anatomic and physiologic integrity of only one fascicle it is reasonable to expect that stress such as anoxemia or drugs which affect atrioventricular conduction may precipitate the development of heart block.

The question therefore arises whether patients with bifascicular block who require surgery should have a prophylactic temporary pacemaker inserted prior to operation. One also wonders about the safety of administering drugs which influence atrio-

Table I Outline of study†

| Underwent surgery | At risk pacing | Received antiarrhythmic drugs |
|------------------------------|---------------------------|-------------------------------|
| 4 patients—had 38 operations | 10—4 of these had surgery | 17—3 of these had surgery |

*Total number of patients = 39

‡Age range = 46 to 90 years.

ventricular conduction such as digoxin or quinidine to patients with bifascicular block. The purpose of this paper is to report our experience in 39 patients with bifascicular block who either underwent surgery and/or received antiarrhythmic drugs.

The introduction of His bundle recording⁷ and atrial pacing⁸ provides a clinical method of studying the electrophysiologic mechanism of atrioventricular conduction

From the Division of Cardiology Department of Medicine and the Department of Anesthesiology New York Medical College, Metropolitan Hospital Center, New York, N.Y.

This study was supported by a Grant in Aid from the Dutchess County Heart Association.

Received for publication Oct. 10, 1972.

Reprint requests to Yusuf Z. Yuceoglu, MD, New York Medical College—Metropolitan Hospital Center, First Avenue, 87th St., New York, N.Y. 10019.

MD Good Fellow in Cardiology

Table 11 Clinical and electrocardiographic features of patients who underwent surgery

| Etiology of heart disease | Number | Sex | | Syncope | Congestive cardiac failure | Cardiomegaly | Functional disturbance | | | |
|-------------------------------|--------|------|--------|---------|----------------------------|--------------|------------------------|----|-----|----|
| | | Male | Female | | | | I | II | III | IV |
| Atherosclerotic heart disease | 9 | 6 | 3 | | 2 | 7 | 5 | 2 | 2 | |
| Hypertensive heart disease | 3 | 3 | — | — | — | 2 | 2 | 1 | — | |
| Etiology not known | 12* | 7 | 5 | — | — | — | 12 | — | — | |
| Total | 24 | 16 | 8 | 1 | 2 | 9 | 19 | 3 | 2 | |
| | | 24 | | | | | 24 | | | |

*Except one patient in whom the etiology may have been traumatic

†New York Heart Association¹⁰

‡Abbreviations and symbols: I = first degree AV block; LBBB = left bundle branch block; RBBB = right bundle branch block

These techniques were applied to ten patients in order to determine whether atrial pacing might be predictive of the development of heart block in persons with bifascicular block.

Materials and methods

The 39 patients with bifascicular block forming the basis of this report were referred to members of the Cardiology Staff at New York Medical College (Lower and Fifth Avenue Hospital and Metropolitan Hospital) for various reasons during the period of June, 1968 to June, 1972. The charts of these patients were reviewed for a history of syncope, evidence of cardiovascular disease, or related conditions such as diabetes mellitus. The criteria for bifascicular block were as follows:

1. Left anterior hemiblock, that is, left axis deviation more negative than -30 degrees (in the absence of inferior wall myocardial infarction or emphysema) and right bundle branch block—i.e., the presence of QRS ≥ 0.12 sec in duration, the presence of wide S waves in Lead I and V_4 and wide R waves in Leads V_1 and V_6 .¹⁰

2. Left posterior hemiblock—i.e., right axis deviation of $\geq +90$ degrees (provided right ventricular hypertrophy, pulmonary emphysema, vertical heart due to slender body build, and extensive lateral wall myo-

cardial infarction could be excluded) and right bundle branch block.¹⁰

3. Left bundle branch block (that is, a QRS of ≥ 0.12 sec in duration, the absence of septal Q waves, the presence of notching of the QRS in the left sided precordial leads and ST T waves in the direction opposite to that of the QRS in the left sided leads)¹⁰ and PR interval greater than 0.20 sec. Patients undergoing surgery were selected for insertion of prophylactic temporary pacemakers at random provided the patient consented to the procedure.

The pacemakers were introduced percutaneously into a femoral vein or by cut down into an antecubital vein under local anesthesia.

The tip of the catheter was positioned in the apex of the right ventricle under fluoroscopic control. The proximal terminals were connected to a Medtronic demand pacemaker. The pacemaker was left in position on demand mode until several days postoperatively.

The electrocardiogram (ECG) of all patients was monitored on an oscilloscope throughout surgery and in the recovery room.

The presence of any arrhythmia, conduction delay, or triggering of the pacemaker was recorded. Complications such as hypotension and hypoxemia were noted. Com-

31 m 86
h mb 2

| Type of surgery | | Type of anaesthesia | | | Type of block | | Electrocardiogram | | |
|-----------------|-------|---------------------|--------|----------|---------------|------------|--------------------|----------------------------|--------------|
| Major | Minor | General | Spinal | Regional | LAH & RBBB | LBBB + LAH | PR interval (msec) | Mean QRS axis (LAH + RBBB) | QRS duration |
| 2 | 4 | 2 | 11 | 3 | 9 | | 0.11 to 0.24 | -45 to -90 | 0.12 to 0.16 |
| 1 | 3 | 1 | - | 2 | 3 | - | 0.16 | -30 to -60 | 0.12 |
| 1 | 8 | 10 | 7 | 2 | 9 | 3 | 0.12 to 0.18 | -30 to -90 | 0.12 to 0.16 |
| 13 | 15 | 13 | 18 | 7 | 21 | 3 | | | |
| 38 | | 38 | | | 24 | | | | |

1 = left anterior hemiblock

plications from the insertion of the temporary pacemaker were also recorded.

His bundle recordings and atrial pacing was performed using the method described by Damato and co-workers.⁸ The A-H and H-V intervals were measured at each atrial pacing rate and the rate at which heart block developed and the site of the block were noted.

Results

Of 39 patients who were studied, 24 had various surgical procedures, 12 received antiarrhythmic drugs, and 10 had atrial pacing (Table I).

There were 27 men and 12 women. Their ages ranged from 46 to 90 years with a mean of 68 years. Tables II, III, and IV list their pertinent clinical and electrocardiographic features. There were 12 patients with atherosclerotic heart disease as evidenced by myocardial infarction or angina pectoris, eight with probable atherosclerotic heart disease (but without a definite history of angina pectoris or myocardial infarction), and six patients with hypertensive cardiovascular disease. Eleven patients out of the 39 had no evidence of heart disease other than the conduction abnormality. One patient had aortic incompetence due to chronic rheumatic heart disease, and one patient presented with

bifascicular block following an automobile accident in which he suffered trauma to his chest.

Surgical group. The 24 patients underwent 38 operations. Six patients underwent more than one operation and one patient had six operations during the period under study. Of the 38 surgical procedures, 23 were classified as major and 15 as minor. Laparotomy, prostatectomy, hysterectomy, and cataract extraction were classified as minor.

Twelve of the operations were performed under general anaesthesia, 17 under spinal, and 7 under regional. The general anaesthetic agent used was either halothane or nitrous oxide.

Of the 38 operations, 13 were performed with the prior insertion of a prophylactic pacemaker, 9 of these being major and 4 minor operations. Fourteen major and 11 minor operations were performed without the prior insertion of a pacemaker (Table V).

There were no complications due to the insertion of the temporary pacemaker in any patient.

None of the 24 patients developed either transient or permanent heart block of any degree during surgery or during the first postoperative week. Two patients devel-

Table IV Clinical and electrocardiographic features of patients who had atrial pacing

| Case no | Age | Sex | Etiology of heart disease | Syncope | CHF | Cardiomegaly | Functional classification | ECG—P R interval (sec) | Mean QRS axis | QRS duration (sec) | Had surgery |
|---------|-----|-----|---------------------------|---------|-----|--------------|---------------------------|------------------------|---------------|--------------------|-------------|
| 1 | 64 | F | ASHD | — | — | Yes | II | 0 16 | -35 | 0 14 | Yes |
| 2 | 88 | M | ASHD | — | — | — | II | 0 20 | -60 | 0 16 | — |
| 3 | 62 | F | HHB | — | — | Yes | II | 0 20 | -60 | 0 12 | — |
| 4 | 70 | M | ASHD | Yes | Yes | Yes | III | 0 20 | -60 | 0 14 | Yes |
| 5 | 70 | F | HHB | — | — | Yes | III | 0 20 | -60 | 0 17 | — |
| 6 | 72 | M | ASHD | — | Yes | Yes | III | 0 30 | -45 | 0 16 | — |
| 7 | 74 | M | ASHD | — | Yes | Yes | II | 0 24 | -60 | 0 12 | — |
| 8 | 65 | M | Etiology not known | — | — | — | I | 0 24 | -60 | 0 12 | Yes |
| 9 | 46 | M | RHD AI | — | Yes | Yes | III | 0 24 | N | 0 12 | — |
| 10 | 70 | M | ASHD | — | — | Yes | II | 0 24 | N | 0 18 | Yes |

*Abbreviations: ECG = electrocardiogram N = normal

Table V Number of operations performed with and without prophylactic pacemakers

| | No of operations | No of patients |
|--------------------------------|------------------|----------------|
| With prophylactic pacemaker | 13 | 11 |
| Without prophylactic pacemaker | 25 | 13 |
| Total | 38 | 24 |

operative patient with presumed bifascicular block is not warranted. Our findings in 24 patients who underwent a total of 38 surgical procedures, 23 of which were major, are similar. One patient had trifascicular block as demonstrated by the His bundle study. None of the patients developed transient or permanent heart block during surgery or during the first postoperative week. Although the experience of Berg and Kotler¹⁴ and ours involves only 54 patients undergoing a total of 74 surgical procedures, it appears that under the controlled conditions of modern anesthesia and surgery the risk of developing heart block in patients with bifascicular block is small. It may be that release of catecholamines during surgery in fact facilitates atrioven-

tricular conduction. Isoproterenol has been shown to shorten the A-H interval at any given heart rate.⁸ The need for the routine insertion of a temporary cardiac pacemaker prior to surgery has not been demonstrated. It is to be noted, however, that there were no patients with left posterior hemiblock and right bundle branch block who underwent surgery in either Berg and Kotler's¹⁴ series or ours.

Patients with this particular type of bifascicular block may have a different propensity to development of heart block compared to the other types observed.⁴ There is evidence that in patients with posterior hemiblock the disease process is more extensive and more diffuse.¹⁵ All one of our patients had thoracotomies. It is possible that patients with bifascicular block undergoing chest surgery may be more prone to develop heart block due to the action of various cardiovascular reflexes.

It is well known that digitalis does not cause impairment of intraventricular conduction. However, digoxin does prolong conduction in the atrioventricular node⁸ and disease of the His-Purkinje system is not infrequently associated with delayed conduction through the A-V node.¹⁶

In our own cases four of eight patients with bifascicular block also had prolonged AV conduction. Quinidine and procaine

Table VI Results of His bundle studies and atrial pacing in ten patients with bifascicular block

| Electrocardiogram | Case no | Basal state | | | Atrial pacing | | | Block | | |
|--|---------|-------------|-----------|-----------|---------------|-----------|-----------|-------|------------|-----------------|
| | | HR* | AH (msec) | HV (msec) | Rate | AH (msec) | HV (msec) | Pace | Type | Site |
| Abnormal LAD + RBBB + normal PR | 1 | 170 | 160 | 40 | 130-165 | 160-230 | 40-50 | 165 | Wenckebach | Proximal to His |
| | 2 | 104 | 90 | 45 | 170-180 | 90-170 | 45-60 | 140 | None | None |
| | 3 | ■ | 90 | 0 | — | — | — | — | — | — |
| | 4 | 7 | 125 | 30 | — | — | — | 140 | Wenckebach | — |
| Abnormal LAD + RBBB + prolonged PR | 5† | — | 110 | — | — | — | — | 140 | Wenckebach | — |
| | 6 | 83 | 110 | 140 | 90-115 | 100-110 | Constant | 115 | Nobitz II | Distal to His |
| | 7 | 79 | 165 | 80 | 90-145 | 165-30 | Constant | 145 | Wenckebach | Proximal to His |
| | 8 | 45 | 100 | 00 | — | — | — | — | — | — |
| Complete LBBB + prolonged PR | 9 | 10 | 165 | 40 | 110-160 | 165-70 | Constant | 160 | Wenckebach | Proximal to His |
| | 10 | 81 | 150 | 60 | 95-105 | 150-700 | Constant | 100 | Wenckebach | Proximal to His |

Atrial pa. = Atrial pacing; HV = HV interval; HR = heart rate; ■ = data not done because of technical difficulties; HR = heart rate.

may do affect the intraventricular conduction and as shown by Damato and colleagues¹⁸ prolong the HV interval on His bundle recording. One might therefore expect that in patients with bifascicular block any of these drugs may further prolong conduction and possibly induce heart block. We have data on only 12 patients in none of whom was there an exacerbation of conduction impairment by the use of these drugs in average therapeutic doses. The number of patients studied however is too small to make a definitive statement.

It is self-evident that it would be useful to have a method for identifying those individuals with bifascicular block who are prone to develop heart block as part of their natural course. This would allow the selection of patients for prophylactic pacemakers in order to avoid a possible fatal Stokes Adams attack.

His bundle recordings allow more accurate localization of the conduction defect than does the electrocardiogram. Thus even in the presence of a normal PR interval A-V nodal conduction may be delayed. This was found in one of our cases (Case 1). On the other hand a prolonged PR interval in the presence of bundle branch block

may be due to disease in the AV node rather than in the contralateral bundle (Case 9).

If there is impairment of conduction in two fascicles only the HV time would be expected to be normal.¹⁹ When it is prolonged as was the finding in 5 of our 9 patients (Cases 3, 6, 7, 8 and 10) more diffuse disease involving either the His bundle or the peripheral Purkinje system can be postulated.

Atrial pacing has also been used to study atrioventricular conduction. It is known that individuals without evidence of any conduction defect develop heart block on atrial pacing at high rates.²⁰ Haft and associates¹⁸ studied His bundle recordings during atrial pacing in 14 patients with bifascicular block and left anterior hemiblock and in four with right bundle branch block and right axis deviation. In all these patients the HV interval remained constant and Wenckebach periods developed proximal to the His bundle as in normals. Cheng²¹ studied ten cases with right bundle branch block and left axis deviation by atrial pacing. Nine patients behaved normally as described above and none of these patients developed complete heart block for a follow up period of up to 30 months.

One patient developed transient complete heart block during pacing at the rate of 120 beats per minute the block being distal to the His bundle. This patient developed Mobitz Type II block and complete heart block one year later.

Berkowitz and associates¹⁹ studied six patients with right bundle branch block and left anterior hemiblock. One of these patients developed Mobitz Type II block distal to His bundle on atrial pacing and this patient subsequently developed complete heart block. In the cases reported here, two out of seven patients studied developed heart block at rates below those found in normals. Only a follow up period will show whether these patients will develop heart block and whether this will occur earlier as compared to the other patients in the group.

In conclusion, our findings indicate that patients with bifascicular block undergoing surgery do not require the routine insertion of cardiac pacemakers prior to operation. The method of His bundle recording during atrial pacing deserves further study as a possible predictive tool in patients with bifascicular block.

Summary

The effect of surgery, arrhythmic drugs and atrial pacing on the atrioventricular conducting system was studied in 39 patients with bifascicular block.

Twenty nine patients underwent 38 operations, 23 of which were major. A temporary cardiac pacemaker was inserted prior to surgery in 13 cases. The remaining 25 operations were performed without prophylactic pacemakers. In no patient did heart block develop either during surgery or in the first postoperative week.

Twelve patients with bifascicular block received digoxin and either quinidine or procainamide in the usual doses. In none was exacerbation of the atrioventricular conduction defect observed.

Ten patients had His bundle recordings during atrial pacing performed at gradually increasing rates until second degree heart block developed. Only two patients developed atrioventricular block at pacing rates below normal, one proximal and one distal to the His bundle.

The authors wish to thank Dr. Jonas Beregov for referring several patients and Drs. Joel Blum and Ralph Schneebaum for help in some of the procedures.

REFERENCES

1. Rosenbaum M. B. and Lepeschkin E. Bilateral bundle branch block. *Am Heart J* 50: 1955.
2. Lenègre J. Etiology and pathology of bilateral bundle branch block in relation to complete heart block. *Progr Cardiovasc Dis* 6: 409-15.
3. Lepeschkin E. The electrocardiographic diagnosis of bilateral bundle branch block in relation to heart block. *Progr Cardiovasc Dis* 6: 1964.
4. Rosenbaum M. B. The hemiblocks: diagnostic criteria and clinical significance. *Mod Concepts Cardiovasc Dis* 34: 141-1970.
5. Castellanos A. Jr. and Leberer, L. Diagnosis of isolated and combined block in the bundle branches and the divisions of the left branch. *Circulation* 48: 971-1971.
6. Lasser R. P., Haft J. I. and Friedberg C. Relationship of right bundle branch block to marked left axis deviation to complete heart block and syncope. *Circulation* 37: 429-1968.
7. Scherlag B. J., Lau S. H., Helfant R. H. et al. Catheter technique for recording His bundle activity in man. *Circulation* 39: 13-1969.
8. Damato A. N., Liu S. H., Helfant R. H. et al. Study of atrioventricular conduction in man using electrode catheter recording of His bundle activity. *Circulation* 39: 187-1969.
9. Kulbertus H. and Collignon P. Association of right bundle branch block with left superior or inferior intraventricular block: its relation to complete heart block and Adams Stokes syndrome. *Br Heart J* 31: 435-1969.
10. Diseases of the heart and blood vessels. Nomenclature and criteria for diagnosis. Criteria Committee of the New York Heart Association. Boston 1969. Little Brown & Company. p. 421-424.
11. Rosen K. M. The contribution of His bundle recording to the understanding of cardiac conduction in man. *Circulation* 43: 961-1971.
12. Scanlon P. J., Pryor R. and Blount S. Right bundle branch block associated with left superior or inferior intraventricular block. *Circulation* 42: 1123-1970.
13. Cheng T. O. Atrial pacing: its diagnostic and therapeutic applications. *Progr Cardiovasc Dis* 14: 230-1971.
14. Berg R. and Kotler M. N. The significance of bilateral bundle branch block in the preoperative patient. *Chest* 59: 62-1972.
15. Narula S. and Samet P. Right bundle branch block with normal left or right axis deviation. *Am J Med* 51: 432-1971.
16. Rosen K. M., Lau S. H. and Damato A. N. Effect of lidocaine on atrioventricular and intraventricular conduction in man. *Am J Cardiol* 25: 1-1970.
17. Schulenburg R. M. and Durrer D. Observa-

- tions on atrioventricular conduction in patients with bilateral bundle branch block. *Circulation* 41:967 1970
- 18 Haft J I, Weinstock M, De Guz R et al. Assessment of atrioventricular conduction in left and right bundle branch block using His bundle electrogram and atrial pacing. *Am J Cardiol* 27:474 1971
- 19 Berkowitz W D, Lau S H, Patton R D, Rosen R M and Dimato A V. The use of His bundle recordings in the analysis of unilateral and bilateral bundle branch block. *Am Heart J* 81:340 1971

One patient developed transient complete heart block during pacing at the rate of 120 beats per minute, the block being distal to the His bundle. This patient developed Mobitz Type II block and complete heart block one year later.

Berkowitz and associates¹⁹ studied six patients with right bundle branch block and left anterior hemiblock. One of these patients developed Mobitz Type II block distal to His bundle on atrial pacing and this patient subsequently developed complete heart block. In the cases reported here two out of seven patients studied developed heart block at rates below those found in normals. Only a follow up period will show whether these patients will develop heart block and whether this will occur earlier as compared to the other patients in the group.

In conclusion our findings indicate that patients with bifascicular block undergoing surgery do not require the routine insertion of cardiac pacemakers prior to operation. The method of His bundle recording during atrial pacing deserves further study as a possible predictive tool in patients with bifascicular block.

Summary

The effect of surgery, antiarrhythmic drugs, and atrial pacing on the atrioventricular conducting system was studied in 39 patients with bifascicular block.

Twenty nine patients underwent 38 operations, 23 of which were major. A temporary cardiac pacemaker was inserted prior to surgery in 13 cases. The remaining 25 operations were performed without prophylactic pacemakers. In no patient did heart block develop either during surgery or in the first postoperative week.

Twelve patients with bifascicular block received digoxin and either quinidine or procainamide in the usual doses. In none was exacerbation of the atrioventricular conduction defect observed.

Ten patients had His bundle recordings during atrial pacing performed at gradually increasing rates until second degree heart block developed. Only two patients developed atrioventricular block at pacing rates below normal, one proximal and one distal to the His bundle.

The authors wish to thank Dr. Jonas Beregovich for referring several patients and Drs. Joel Dugash and Ralph Schneebaum for helping in some of the procedures.

REFERENCES

1. Rosenbaum M B and Lepschkin E Bilateral bundle branch block. *AM HEART J* 50:38 1955
2. Lemire J Etiology and pathology of bilateral bundle branch block in relation to complete heart block. *Progr Cardiovasc Dis* 6:409 1964
3. Lepschkin E The electrocardiographic diagnosis of bilateral bundle branch block in relation to heart block. *Progr Cardiovasc Dis* 6:445 1964
4. Rosenbaum M B The hemiblocks: diagnostic criteria and clinical significance. *Mod Concepts Cardiovasc Dis* 31:141 1970
5. Castellanos A Jr and Lemberg L Diagnosis of isolated and combined block in the bundle branches and the divisions of the left branch. *Circulation* 48:971 1971
6. Lasser R P Haft J I and Friedberg C H Relationship of right bundle branch block and marked left axis deviation to complete heart block and syncope. *Circulation* 47:429 1968
7. Scherlag B J, Lau S H, Helfant R H et al Catheter technique for recording His bundle activity in man. *Circulation* 39:13 1969
8. Damato A N, Lau S H, Helfant R H et al Study of atrioventricular conduction in man using electrode catheter recording of His bundle activity. *Circulation* 39:287 1969
9. Kuibertus H and Collignon P Association of right bundle branch block with left superior or inferior intraventricular block: its relation to complete heart block and Adams Stokes syndrome. *Br Heart J* 31:435 1969
10. Diseases of the heart and blood vessels. Nomenclature and criteria for diagnosis. Criteria Committee of the New York Heart Association. Boston 1969 Little Brown & Company p 421-424
11. Rosen K M The contribution of His bundle recording to the understanding of cardiac conduction in man. *Circulation* 43:961 1971
12. Scanlon P J, Pryor R and Blount S Right bundle branch block associated with left superior or inferior intraventricular block. *Circulation* 42:1123 1970
13. Cheng T O Atrial pacing: its diagnostic and therapeutic applications. *Progr Cardiovasc Dis* 14:230 1971
14. Berg R and Kotler M N The significance of bilateral bundle branch block in the preoperative patient. *Chest* 59:62 1972
15. Narula S and Samet P Right bundle branch block with normal left or right axis deviation. *Am J Med* 51:432 1971
16. Rosen K M, Lau S H and Damato A N Effect of lidocaine on atrioventricular and intraventricular conduction in man. *Am J Cardiol* 25:1 1970
17. Schulenburg R M and Durrer D Observa-

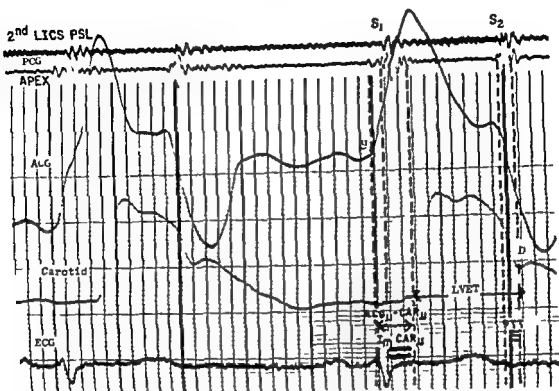


Fig 1 Subject young woman Simultaneous recording from top to bottom of PCG = phonocardiogram (second left intercostal space parasternal line) APEX = apex phonocardiogram ACG = apexcardiogram Carotid = external carotid pulse ECG = electrocardiogram S_1 and S_2 = First and second heart sounds D = diastolic notch u = upstroke of carotid and apexcardiogram $ACG-CAR$ = isovolumic contraction time from upstroke of ACG to upstroke of carotid pulse I_m-CAR = isovolumic contraction time from high frequency component of first heart sound to upstroke of carotid PTT = pulse transmission time Paper speed = 100 mm per second

splitter to divert it to another AC amplifier with the band pass filter set at 0.1 Hz to 15 Hz. The external right carotid arterial pulse tracings were recorded by means of a funnel shaped pickup attached to a Sanborn pulse wave transducer 210s1D.

Isovolumic contract time (IVCT) was determined by the two methods described by Spodick and Kumar.⁶

1 Onset of the first rapid vibrations of the first heart sound to the upstroke of the carotid pulse (I_m-CAR). This was also corrected for the pulse transmission time (I_m-CAR minus PTT) (see Fig 1).

2 Onset of the apexcardiogram to the upstroke of the carotid pulse ($ACG-CAR$). This was also corrected for pulse transmission time ($ACG-CAR$ minus PTT) (see Fig 1). Because indirect recordings cannot be made from the

ascending aorta the onset of pressure rise in the carotid artery was utilized instead. The time for the transmission of the pulse wave from aorta to the carotid is designated as the pulse transmission time. The aortic second sound is coincident with the aortic incisure and therefore the pulse transmission time was measured from the aortic second sound to the carotid incisure. This was deducted from the time of onset of the carotid upstroke to indicate beginning of ejection and endpoint of isovolumic contraction time.

The left ventricular ejection time (LVET) was obtained from the beginning of the upstroke to the trough of the incisura of the carotid pulse tracing. The R-R interval was used to determine the heart rate. The left ventricular ejection time was corrected by the formula outlined by Weissler

Systolic time intervals in pregnancy and the postpartum period

Shirley Rubler, M D, F A C C

Ralph Schneebaum, M D†

Nina Hammer, B A

New York N Y

Noninvasive techniques have recently been employed to evaluate myocardial function^{1,6}. The cardiac performance has been assessed after exercise⁷ during sustained hand grip after the administration of such agents as amyl nitrate and phenylephrine, and during isorhythmic dissociation⁸.

This investigation was designed to assess the effect of a sustained expansion of blood volume on myocardial contractility in normal subjects. The isovolumic contraction time (IVCT) and the left ventricular ejection time index (LVETI) were studied in pregnant and postpartum subjects. Furthermore the influence of the assumption of the supine and upright positions upon these responses were evaluated.

Materials and methods

Sixteen normal women 18 to 36 years of age were studied at the time of maximal expansion of blood volume (at the twenty-seventh to thirty-third weeks of gestation). Six of the original group were re-evaluated six weeks postpartum. In addition the ten subjects who did not return for re-study were replaced by a comparable age-matched group of postpartum women. The

findings for the original six patients on whom the investigation was repeated in the postpartum period were similar to those obtained for the ten substituted postpartum subjects. Both sets were therefore combined for the final evaluation of the entire series (See under Results).

All subjects were referred for evaluation from the obstetrical and family planning clinics of a large metropolitan hospital and were entirely free of any medical disability or obstetrical complication.

Simultaneous external phonocardiograms, electrocardiograms, right carotid tracings and apexcardiograms were recorded at rest in the supine position and after assuming the sitting position.

The recordings were obtained at a paper speed of 100 mm per second on an eight-channel recorder (Sanborn Hewlett Packard 4568 A). Phonocardiograms were obtained at the apex with a 31050 A/B contact sensor (Sanborn Hewlett Packard) using frequency responses 50 to 200 Hz. Another microphone was placed at the second left intercostal space at the parasternal line. The apexcardiogram was procured from the same contact sensor used to record heart sounds by means of a signal

From the New York Medical College Metropolitan Hospital Center New York N Y
Received for publication Oct 10 1972

Reprint requests to Shirley Rubler M D Metropolitan Hospital Center 1901 First Ave New York N Y 10029
†Dr Schneebaum died since this paper was written

August 1973 Vol

positions

| LVETI | | SD | HRR | |
|----------------|------|------|-----------------|------|
| (Mean in msec) | | | (Beats per min) | |
| 425.1 | (16) | 13.5 | 87.2 | (16) |
| 412.8 | (16) | 14.2 | 94.3 | (16) |
| 408.6 | (6) | 17.0 | 70.7 | (6) |
| 406.1 | (10) | 18.5 | 71.9 | (10) |
| 407.0 | (16) | 17.4 | 71.4 | (16) |
| 393.4 | (6) | 6.4 | 74.8 | (6) |
| 397.1 | (10) | 14.2 | 85.4 | (10) |
| 397.9 | (16) | 11.6 | 81.6 | (16) |

| LVETI | |
|---------|--|
| 0.001 | |
| < 0.001 | |
| < 0.01 | |
| < 0.001 | |

SD = standard deviation; I_m CAR = stroke interval; PTT = pulse transmission time; LVETI = left ventricular ejection time; LVEI = left ventricular ejection index

applicable to the pregnant woman and therefore minimized the expected change in isovolumic contraction time upon the assumption of the sitting position. However, this explanation does not appear to be feasible because the left ventricular ejection time shortened in the pregnant subject in the upright position while in the cardiac patient with increased blood volume the ventricular ejection time remained unchanged.

The left ventricular ejection time index (LVETI) in the antepartum subjects both in the supine and upright positions was longer than the comparable LVETI for that position in the postpartum period. This again could be attributed to the expanded blood volume and increased stroke volume of pregnancy which operated in both positions. In the erect posture however the LVETI shortened significantly

and to the same degree in the antepartum as in the postpartum period.

It would seem therefore that when the blood volume is at its peak level during the gestation period and the myocardial fibers are lengthened the alteration in systolic time intervals reflects these physiological changes. The isovolumic contraction time is abbreviated in response to the greater initial fiber length, the left ventricular ejection time is lengthened due to the increase in stroke volume. Pregnant women (at least until the terminal weeks of pregnancy) respond to changes in position from the supine to the upright in the expected manner and no evidence of uterine compression upon inferior vena caval flow was noted.

The data presented here for the normal pregnant woman could be used in assessing the functional status of cardiac patients in the antepartum and early postpartum periods and for establishing prognostic and therapeutic criteria in the future for these patients.

Summary

Systolic time intervals in the supine and upright positions were obtained in 16 antepartum subjects at maximal expansion of blood volume (27 to 33 weeks) and six weeks postpartum. Isovolumic contraction time (IVCT) was determined by 2 methods: (1) first heart sound to carotid upstroke interval (I_m CAR) and minus pulse transmission time (I_m CAR - PTT) and by (2) upstroke of apexcardiogram to upstroke of carotid (ACG CAR) and minus pulse transmission time (ACG CAR - PTT). The IVCT was short in the antepartum supine subject (I_m CAR was 60.5 ± 4.3 msec) and 31.7 msec when corrected for PTT) and became longer in the upright position (63.7 msec [SD 6.1] and 35.6 msec respectively).

In the postpartum state the supine IVCT was significantly longer than in the antepartum state (72.2 msec [SD 9.3] and 42.2 msec when corrected for PTT) (P < 0.001) and became longer in the sitting postpartum subject (85.4 msec [SD 8.8] and 56.1 msec respectively).

The left ventricular ejection time index was 42.51 msec (SD 13.5) in the antepartum supine subject and demonstrated

Table 1 Systolic time intervals in pregnancy and the postpartum period in supine and sitting

| Group | Im CAR † | | SD | Im CAR PTT | | ACG CAR | | SD | ACG CAR PTT | |
|-----------------------|----------------|-------|------|----------------|------|---------|----------------|------|-------------|----------------|
| | (Mean in msec) | | | (Mean in msec) | | | (Mean in msec) | | | (Mean in msec) |
| I Antepartum lying | 60.5 | (16)§ | 4.3 | 31.7 | (16) | 91.0 | (9) | 7.8 | 63.6 | (9) |
| II Antepartum sitting | 63.7 | (16) | 6.1 | 35.6 | (16) | 95.0 | (6) | 5.9 | 66.5 | (9) |
| III Postpartum lying | | | | | | | | | | |
| *Group A | 70.7 | (6) | 11.2 | 40.2 | (6) | 100.3 | (5) | 15.9 | 69.9 | (5) |
| †Group B | 73.2 | (10) | 8.5 | 43.2 | (10) | 104.2 | (6) | 11.7 | 74.1 | (6) |
| Total cries | 72.2 | (16) | 9.3 | 42.2 | (16) | 107.4 | (11) | 13.1 | 72.2 | (11) |
| IV Postpartum sitting | | | | | | | | | | |
| *Group A | 85.0 | (6) | 8.6 | 56.9 | (6) | 112.5 | (5) | 9.6 | 84.4 | (5) |
| †Group B | 85.6 | (10) | 9.4 | 56.6 | (10) | 114.4 | (7) | 10.6 | 85.4 | (7) |
| Total series | 85.4 | (16) | 8.8 | 56.7 | (16) | 113.6 | (12) | 9.8 | 84.9 | (17) |

| P values | Im CAR | ACG CAR |
|--|---------|------------|
| Antepartum lying vs sitting | < 0.02 | NS (> 0.1) |
| Postpartum lying vs sitting | < 0.001 | < 0.01 |
| Antepartum lying vs postpartum lying | < 0.001 | < 0.07 |
| Antepartum sitting vs postpartum sitting | < 0.001 | < 0.001 |

*Group of original patients that returned postpartum for restudy

†Group of patients substituted for defectors

§Abbreviations: Im CAR = isovolumic contraction time from high frequency component of first heart sound to upstroke of carotid pulse; PTT = pulse transmission time; ACG CAR = ACG carotid pulse transmission time; ACG CAR PTT = isovolumic contraction time from upstroke of apexcardiogram to upstroke of carotid pulse; IIR = heart rate; NS = not statistically significant

Numbers in parentheses indicate number of patients

that we observed. Burg and associates¹⁹ obtained similar results when studying subjects in the second trimester of pregnancy. These authors, however, found that the pre-ejection period was longer in the supine position in the third trimester and attributed this to the effect of the enlarged uterus on the venous return. This factor assumed greater significance in their patients than in ours. One reason for the discrepancies in these observations may have been that our patients' systolic time intervals were recorded at the time of maximum increase in blood volume and not in the last weeks of pregnancy. The effect of the gravid uterus upon the venous return in this late antepartum period might be greater than the alterations caused by the expanded blood volume.

A small but significant lengthening of the isovolumic contraction time was noted

when the patients were upright. However, after delivery in the postpartum women the isovolumic contraction time was more significantly lengthened on assuming the sitting position. This is similar to the effect that Stafford and co-workers²⁰ noted in other normal subjects and has been attributed to the decrease in venous return occasioned by the erect posture. In the antepartum women the expanded blood volume may have minimized the effect of the upright position on the IVCT. This response is similar to that observed in patients with congestive heart failure. The latter subjects demonstrated an unresponsiveness to head up tilt and the systolic time intervals were unchanged in this position.²¹ They attributed these findings to the antigravitational effects of hypervolemia, the elevated tissue pressure and increased venous tone. This might be

Experimental and laboratory reports

Mathematical relationship between automaticity of the sinus node and the AV junction

Ferdinand Urthaler M D
Charles R Katholi Ph D
Josiah Macy Jr Ph D
Thomas N James M D
Birmingham Ala

Any attempt to define the sequence of events which regulates the normal heart rate must in essence refer to mechanisms which accelerate or slow the sinus node. There are, however, alternate pacemakers and the major substitute pacemaker of the mammalian heart is in or near the AV (atrioventricular) node-His bundle region.^{1,2} Despite the current uncertainties as to the exact location of this AV junctional pacemaker, it recently became possible to explore selectively the direct and indirect chronotropic responses of this subsidiary pacemaker in the intact canine heart *in situ*.^{3,4}

For the sake of systematic investigation of normal and abnormal cardiac rhythms it is useful to have quantitative information about stable sinus and AV junctional rates and from this to establish whether or not there is any mathematical correlation between these two variables. We describe in this paper the data obtained in 50 dogs and their mathematical analysis.

Method

Fifty mongrel dogs weighing 14 to 23 kilograms were anesthetized with intra-

venous (IV) administration of pentobarbital sodium (30 mg per kilogram of body weight). Ventilation was maintained through a cuffed endotracheal tube with an intermittent positive pressure Harvard pump supplying room air at 14 to 16 strokes per minute and at a tidal volume of 225 ± 25 ml. Each dog was anticoagulated with sodium heparin (2 mg per kilogram of body weight). IV Central aortic and right atrial pressures were measured with catheters positioned by way of the right carotid artery and right external jugular vein and attached to transducers. The chest was opened in the right fourth intercostal space and both the sinus node and the AV node arteries were cannulated with small polyethylene catheters; details of these procedures have been reported.^{5,6}

Throughout each experiment a unipolar wick-electrogram from near the sinus node was recorded on a separate single channel electrocardiograph at 25 mm per second. The same right atrial electrogram was also simultaneously recorded with the two pressures, one or two unipolar electrograms from the right and/or left ventricular surface and a tachogram on a multichannel

From the Cardiovascular Research and Training Center, University of Alabama School of Medicine, Birmingham, Ala. This work was supported in part by grants from the National Heart Institute and the National Institutes of Health (HL 11,310).

Received for publication Sept 5, 1972.

Reprint requests: Thomas N. James, M.D., Cardiovascular Research and Training Center, University of Alabama School of Medicine, Birmingham, Ala. 35294.

corresponding shortening in the erect position (412.8 msec [SD 14.2])

In the postpartum period the LVETI was shorter (407.0 msec [SD 17.4]) and shortest in the upright position (392.9 msec [SD 11.6])

We wish to express our thanks to Miss Seto Chice for her technical assistance and to Dr. Mary Monk and Dr. Marvin Glaser for their help with statistical evaluations. For her assistance in typing we thank Mrs. Leonie Harris.

REFERENCES

1. Jezek V. Clinical value of the polygraphic tracings in the study of the sequence of events during cardiac contraction. *Cardiology* 53:298 1963
2. Weissler A M, Aronow M, Harris N S and White D M. Left ventricular ejection time index in man. *J Appl Physiol* 18:919 1963
3. Isfar F, Cohen I S and Levene H D. The normal apocardiogram. The temporal relationship to electrical, acoustic and mechanical cardiac events. *Circulation* 30:381 1964
4. Kumar S and Spodick D H. Study of the mechanical events of the left ventricle by atrumatic techniques. Comparison of methods of measurement and their significance. *Am Heart J* 80:401 1970
5. Spodick D H and Kumar S. Isovolumetric contraction period of the left ventricle. *Am Heart J* 76:498 1968
6. Weissler A M, Harris N S and Schoenfeld C D. Bedside techniques for the evaluation of ventricular function in man. *Am J Cardiol* 23:577 1967
7. Aronow W S. Isovolumic contraction and left ventricular ejection times. *Am J Cardiol* 26:288 1970
8. Marlen C, Shaver J, Thompson M F, Reddy T S and Leonard J J. Direct correlations of external systolic time intervals with internal indices of left ventricular function in man. *Circulation* 44:419 1971
9. Lees M M, Taylor S H, Scott D B and Kerr M G. A study of cardiac output at rest throughout pregnancy. *J Obstet Gynaecol Br Commonw* 74:319 1967
10. Kerr M G. The mechanical effect of the gravid uterus in late pregnancy. *J Obstet and Gynaecol Br Commonw* 72:513 1965
11. Pritchard J A. Changes in blood volume during pregnancy and delivery. *Anesthesiology* 26:393 1965
12. Hurst J N and Logue R B. The heart. New York 1970. McGraw Hill Book Company, Inc. p 1395
13. Pritchard J A and Adams R H. Erythrocyte production and destruction during pregnancy. *Am J Obstet Gynaecol* 79:751 1960
14. Uelano K and Parer J T. Effects of estrogens on the cardiovascular system of the ewe. *Am J Obstet Gynaecol* 96:400 1966
15. Williams J N. Maternal physiology in pregnancy. In Hellman L M and Pritchard J A, editors. *Obstetrics*. Chapter 8. New York 1971. Appleton Century Crofts Inc.
16. Weissler A M and Schoenfeld C D. Effect of systolic time intervals in heart failure. *Am J Med Sci* 259:4 1970
17. Wallace A G, Mitchell J H, Skinner N S and Siroff S S. Duration of the phases of left ventricular systole. *Circ Res* 12:611 1963
18. Sonnenblick E H. Series elastic and contractile element in heart muscle. Changes in muscle length. *Am J Physiol* 207:1130 1964
19. Burg J I, Dodek I and Kloster F E. Systolic time intervals in pregnancy. (Abstract). *Clin Res* 30:365 1972
20. Stafford R N, Harris W S and Weissler A M. Left ventricular systolic time interval as indices of postural circulatory stress in man. *Circulation* 41:485 1970
21. Ibid

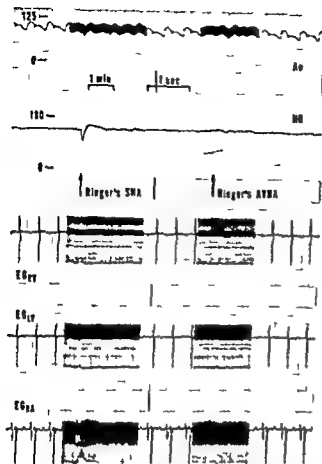


Fig 1 This polygraph illustrates the standard recording technique and chronotropic responses to injections of Ringer's solution into the sinus node artery (SNA) and AV node artery (AVNA) during normal sinus rhythm. Bradycardia during injection and slight post injection tachycardia is characteristic of the response when the nutrient artery perfuses the driving pacemaker, as in the first injection above. There is no discernible effect from an injection into a driven or latent pacemaker (second injection here). Recording channels from above down are aortic pressure (Ao) tachogram (HR) and unipolar electrograms from right ventricular, left ventricular and right atrial surfaces. Pressure is scaled in mm Hg and HR in beats per minute.

polygraph at appropriate speed (usually 0.25 or 25 mm per second). The tachogram was derived from successive R waves of either the right or left ventricular electrogram.

After completion of the initial experimental preparation which includes cannulation of both the sinus node and AV node arteries, control sinus rate was recorded before and after sinus node perfusion with Ringer's solution. It has previously been established that the operative procedures per se have no significant electrophysiologic effects.^{2,6} The sinus rate was considered stable when after injection brady-

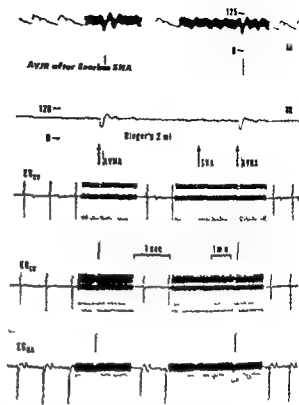


Fig 2 When the AV junction is the driving pacemaker, injection into the AV node artery produces bradycardia but injection into the sinus node artery produces no discernible effect.

cardia and post injection acceleration⁷ the rate remained identical to the one observed prior to injection (Fig 1). To assess the accuracy of the response to selective AV junctional perfusion, 2 ml of acetylcholine chloride (0.1 μ g per milliliter and 1 μ g per milliliter prepared in Ringer's solution), experiments were considered satisfactory only if acetylcholine 0.1 μ g per milliliter produced an immediate complete AV block for 1 second, or if 1 μ g per milliliter similarly blocked AV conduction for at least 2 seconds. To obtain a stable AV junctional rhythm, sinus node activity was gradually depressed by direct perfusion through the sinus node artery with 2 ml of eserine (physostigmine salicylate 10 to 100 μ g per milliliter), validity of this method has been described previously.¹⁴ Such escape AV junctional rhythm persists for the duration of cholinesterase paralysis in the sinus node and in the present experiments averaged 15 ± 5 minutes. AV junctional rate was considered stable when after AV junctional bradycardia and

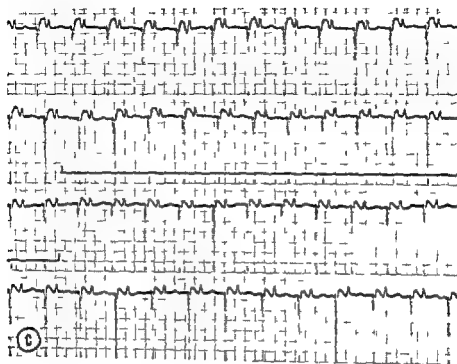


Fig 4C See legend on opposite page

Table I Analysis of variance for regression of AVJR on sinus rate

| Source | df | SS | MS | F test |
|--------------------------------------|----|--------------|--------------|----------|
| Due to slope | 1 | 399 558 4218 | 399 558 4218 | 9 972 31 |
| Additional due to intercept | 1 | 105 09 2 | 105 0972 | 2 62 \S |
| Additional due to lack of linear fit | 16 | 690 3477 | 43 1467 | 1 08 \S |
| Error | 37 | 1 287 1333 | 40 0667 | |
| Total (uncorrected) | 50 | 401 636 0000 | | |

*Significant at $p < 0.001$

Abbreviations: df = degrees of freedom; SS = sum of squares; MS = mean square; \S = significant at $p = 0.10$

that resting AVJR amounts to 66 per cent or two thirds of the resting SR as shown as very close to correct since the calculated value is 64.98 per cent. As with most biologic measurements one must allow for potential sources of experimental error such as those inherent in visual reading of electrograms, variability of recording paper drive speed and the existing slight inherent deviation from linearity of the tachogram in both high and low frequency range. If these and unrecognized other sources of error could be eliminated AVJR would prove to be exactly two thirds of SR.

Although this is speculative. The analysis of variance considering the source due to the slope was significant ($p < 0.001$) whereas considering additional sources due to both the intercept and to the lack of linear fit were not significant ($p = 0.10$). The model $AVJR = \beta SR$ is therefore adequate to explain the relationship between the two variables AVJR and SR. It has further been demonstrated that with a β estimate of the slope being 0.64979 the t test of the slope is significant at a $p < 0.001$ level and that the range of the 95 per cent confidence limits for the mean

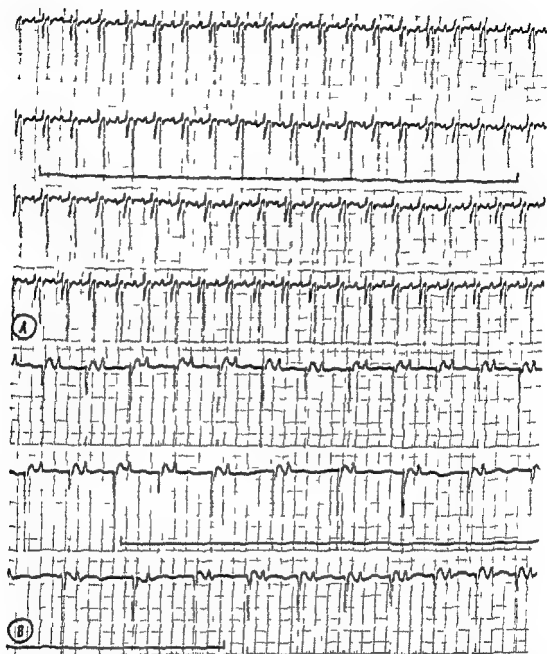


Fig 4 A through C Along with the polygraphs in Figs 1 to 3 the right atrial electrograms (25 mm/sec paper speed) illustrate the usual sequence of experiments reported. Characteristics of the sinus bradycardia during injections into the sinus node artery (SNA) have been published.^{1,2} In A there is no discernible effect from injection of 2 ml Ringer's solution into the AV node artery during normal sinus rhythm. In B a similar injection during AV junctional rhythm produces slowing, but in C an injection into the sinus node artery (after Esarine SNA) has no effect on the AV junctional rhythm. These records are from the same dog illustrated in Figs 1 to 3 and are typical of all other records.

ing between 48 and 126. Fig 6 represents the cumulative distribution of both the sinus and AV junctional rates. As illustrated in Fig 7 by the plot of AV junctional rate (AVJR) versus corresponding sinus rate (SR) each in the same dog the ratio AVJR to SR over the whole range of frequencies is consistently very close to 2/3. The results of statistical treatment³ of the data are summarized in Tables I and II. Under the experimental conditions previ-

ously described the rate of a stable AV junctional escape rhythm amounts to about 66 per cent of the resting control sinus rate.

Discussion

There is a remarkably consistent correlation ($r = 0.942$, $p < 0.001$) for the linear relationship between the rate of stable sinus rhythm and in the same dog the rate of stable AV junctional escape rhythm. A convenient and rem-

SINUS RATE AND ESCAPE AV JUNCTIONAL RATE IN EACH OF 50 DOGS

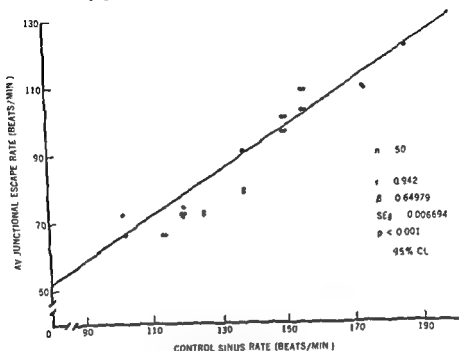


Fig 7 This summary graph illustrates the relationship of escape AV junctional rate to control sinus rate each dot representing the two observations in a given dog of the 50 in the series

minute (mean ± 1 SD) the range being 84 to 192. Following selective eserinization of the sinus node (10 or 100 μ g per milliliter 2 ml) which progressively slows the sinus node until a stable AV junctional rhythm emerges, the mean AV junctional rate was 88 ± 19 beats per minute, ranging between 48 and 126. The plot of AV junctional rates (AVJR) versus corresponding sinus rates (SR) from each dog demonstrates a 2:3 ratio between these two variables. On mathematical analysis of these data there is a consistent correlation ($r = 0.942$, $p < 0.001$) for the linear relationship between AVJR and SR, with a β estimate of the slope being 0.64979. This value is reasonably close to a useful premise that the resting AV junctional rate amounts to 66 per cent or two thirds of the control sinus rate. The model $AVJR = \beta SR$ is adequate to explain the relationship between the two variables AVJR and SR.

REFERENCES

1. Weiskopf J and Eyster J A E. Experiment on the origin and propagation of the impulse in the heart. *Heart* 5:227 1914.
2. Fetter J A E and Weiskopf J. Experiments

on the origin and conduction of the cardiac impulse. VI. Conduction of the excitation from the sino-auricular node to the right auricle and auriculoventricular node. *Arch Intern Med* 18:774 1916.

3. James T N, Bear F S, Frank R J, Lang K F and Tomlinson J C. Selective stimulation suppression or blockade of the atrioventricular node and His bundle. *J Lab Clin Med* 76:740 1970.
4. Lethaler F and James T N. Effects of adenosine and ATP on AV conduction and on AV junctional rhythm. *J Lab Clin Med* 79:96 1972.
5. James T N and Nadeau R A. Direct perfusion of the sinus node. An experimental model for pharmacological and electrophysiological studies of the heart. *Henry Ford Hosp Med J* 10:21 1967.
6. James T N and Hervey E A Jr. Experimental studies on the pathogenesis of atrial arrhythmias in myocardial infarction. *Am Heart J* 63:196 1967.
7. James T N and Nadeau R A. Sinus bradycardia during injections directly into the sinus node artery. *Am J Physiol* 204:9 1963.
8. James T N. Cholinergic mechanisms in the sinus node with particular reference to the action of the nictitatum. *Circ Res* 19:347 1966.
9. Brownlee K A. Statistical theory and methodology in science and engineering. 2nd ed. New York 1965. John Wiley & Sons Inc.

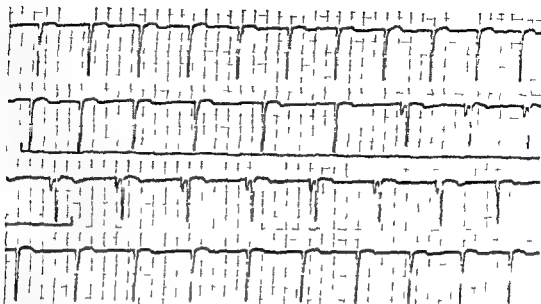


Fig 5 An injection into the AV node artery during AV junctional rhythm (after F-serine SNA) under certain circumstances permit a low escape rate of the exterminized sinus node. See text for discussion.

Table II Final statistics for fit of assumed model $AVJR = \beta SR$

| Variables | Values |
|---|----------|
| Correlation coefficient (r) | 0.947 |
| Variance of points from fitted line (s^2) | 42.3996 |
| Degrees of freedom associated with s^2 | 49.0 |
| Estimate of slope β | 0.64919 |
| Standard error of slope estimate | 0.006694 |
| t test of slope | 97.07* |

*Significant at $p < 0.001$ level

remains narrow over a wide frequency distribution (Fig 7).

It has been common knowledge since the early use of a Stannius ligature in amphibian hearts and from the work of Meek and Eyster^{1,2} on vertebrate hearts that sites of automaticity below the level of the sinus node exhibit decreasing rates of escape activity. We are unaware, however, of any previous systematic analysis of mathematical relations between rates of the sinus node and its major substitute pacemaker observed and documented in vivo. The observation in our experiments that the stable AV junctional rate amounts to about 66 per cent of the control sinus rate corresponds closely to the 67 per cent value which Eyster and Meek predicted for the rate of the canine AV node.

Summary

Stable sinus and AV junctional rates (each in the same animal) were obtained

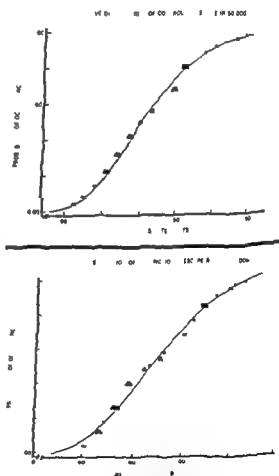


Fig 6 The two graphs illustrate the cumulative distribution of sinus rate and escape AV junctional rate from the 50 dogs in this study.

in 50 open chest dogs anesthetized with pentobarbital. After cannulation of both the sinus node and the AV junctional arteries the control sinus rate was 150 \pm 15 p

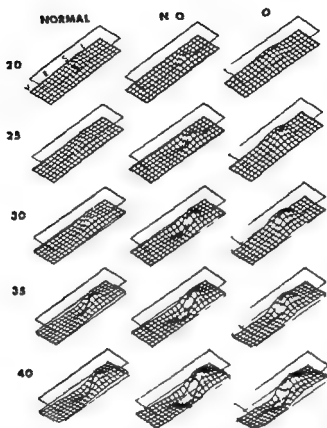


Fig 1 Average isometric projection maps for the group of normal subject, the group with nonobstructive cardiomyopathy (NO) and the group with obstructive cardiomyopathy (O). Landmarks applicable to this and all succeeding figures are indicated in the 20 msec average normal map (upper left). The midsternal line (S), the right midaxillary line (R), the left midaxillary line (L), the vertebral line (V) and the xiphoid process (X) are indicated. The plane above the grid represents the 1 mV level. The numbers in the left hand column indicate the millisecond in ventricular depolarization represented by the surface potential grids from each of the three groups. The anterior bulge most prominent to the left of the sternum in the normal group represents a combination of left to-right septal activation and early activation of the right ventricle.^{1, 2} The mound of positivity increases, begins to move to the left quite obviously by 30 msec, with evidence that right ventricular breakthrough has occurred, demonstrated by the dip of negativity seen at 30 msec. This valley of negativity increases in magnitude and perimeter through 35 and 40 msec as more of the right and left ventricular epicardial surface at their septal attachments, and later more of the free walls become activated. The group with nonobstructive disease shows a somewhat earlier dip of negativity which is increased in magnitude and perimeter over the normal but in general follows the normal sequence of activation. The obstructive group, on the other hand, marked by a more prominent positive bulge which centers slightly more to the right and represents not only septal and normal right ventricular activation but also the influence of those patients in the group who had right ventricular outflow tract obstruction and associated increase in wall thickness of the right ventricle and right aspect of the septum. The mound increases its positive expression and the delayed right ventricular breakthrough occurs at about 35 msec in contrast with earlier breakthrough noted in the other two groups. Electrophysiologic detail is muted by the average maps but the general theme is evident. See text for further discussion.

had biventricular enlargement and seven had predominantly left ventricular enlargement. All had an audible S_3 and S_4 at some time during our evaluation except one who had an S_4 only and one who only had significantly elevated left ventricular end-diastolic pressures and decreased dP/dT associated with left ventricular enlargement but in whom no gallops were heard.

All had a history of a regular intake of alcohol on a daily basis for over five years. All but one had had frank episode of cardiac insufficiency including not only the reported gallops but rales and an initial significant diuretic response to digitalis. No other etiology for their heart failure could be found.

The data were obtained by high fre-

Comparative surface potential patterns in obstructive and nonobstructive cardiomyopathy

Nancy C Flowers M D

Leo G Horan, M D

Augusta Ga

In this study we will present comparative aspects of two hemodynamically different forms of cardiac muscle disease obstructive and nonobstructive both of which cause ventricular enlargement but result in distinctly different expressions of body surface potential. The group of patients with left ventricular outflow obstruction all had nonfamilial idiopathic hypertrophic subaortic stenosis (IHSS). The patients in the remaining group had one of the nonobstructive nonfamilial types of cardiomyopathy.

Methods

A total of 19 patients with cardiomyopathy were studied by means of body surface potential mapping. In every patient the cardiac diagnosis was verified by means of an extensive cardiovascular history and physical examination, standard electrocardiograms (ECG), posteroanterior and lateral roentgenograms, cardiac survey roentgenograms with barium swallow, and vectorcardiography (VCG) loops and high frequency scalar orthogonal leads. In all patients with IHSS and in seven of 13 patients with nonobstructive cardiomyop-

athy, the diagnosis was also supported by cardiac catheterization.

From 142 thoracic sites selected by means of a cylindrical coordinate marking system, QRS complexes were recorded from 30 normal control male subjects, from six male patients with IHSS, and from 13 male patients with nonobstructive cardiomyopathy. The normal subjects ranged in age from 22 to 29 years. Their normal state was verified by the same diagnostic approaches listed above for the patient groups, except cardiac catheterization. In each instance the patients with IHSS had a clearly demonstrated gradient at catheterization from the low left ventricle to the subvalvular chamber, and an increase in gradient with pharmacologic stimulation and after ventricular premature beat. The ages of the patients in the nonobstructive group ranged from 27 to 31 years with a mean of 40. In the obstructive group the age range was 18 to 35 years with a mean age of 41.3. All patients in the nonobstructive category were also completely free of evidence of alternative or combined etiologic possibilities to explain clear cut manifestations of ventricular dysfunction. Six

From the Section of Cardiology, Medical Service, FHD Veterans Administration Hospital and the Department of Medicine, Medical College of Georgia, Augusta, Ga.

This study was supported by an award from the Veterans Administration, United States Public Health Services Grant No. HE 11667, and Grant No. 69-783 from the American Heart Association.

Received for publication Sept. 22, 1972.

Reprint requests to Nancy C. Flowers, M.D., Chief, Section of Cardiology, FHD Veterans Administration Hospital, Augusta, Ga. 30904.

quency techniques fed into an analog to digital converter and ultimately stored at sampling rates of 1 000 bits per second for final processing by means of a PDP 9 computer. These data were then available for display on a storage oscilloscope for hard copy photography in the form of high frequency ECGs, VCG scalar leads or loops¹ and isometric projection maps at 5 msec intervals throughout the QRS complex.

Maps representing the group average were constructed by taking each individual's digitized data throughout his QRS complex obtained from the 142 sampling sites and averaging with the corresponding instant in the QRS of each member of the group.

In Fig. 1 the isometric projection grid for demonstrating changes in body surface potential distribution is illustrated. Land marks which apply to all successive grids are seen in the first normal projection map and apply to each successive map: the mid-teral line (S), the left midaxillary line (L), the right midaxillary line (R), the paravertebral line (V) and the xiphoid process (X). The plane inscribed above each grid represents the 1 mV level. Chest display of positive potential is represented by an outward bulge in the map while negative potential is represented by sinks in the map.

Results

Although the technique of map averaging of several individuals tends to mute electrophysiologic detail it does serve the purpose of presenting in over all theme of group differences. At 20 msec there is an anterior bulge representing transeptal left to right activation which is also contributed to by early activation of the right ventricular wall near to the attachment of the papillary muscle. From about 10 msec on which is not illustrated here there has been some excitation of the right ventricular septal surface.² The net effect of these wave forms on a body surface potential map then is an outward mound to the right of the sternum with a larger outward mound beginning to move slightly anteriorly and leftward as first the septal attachment and later the free wall of the left ventricle becomes activated. At about

30 msec a diffuse elevation occurs which as peak approaches the left midclavicular line at about the fourth intercostal space. Breakthrough of the wave of activation to the right ventricular surface (which is represented somewhat earlier in individual maps) probably accounts for the dip of negativity seen by 30 msec which becomes quite prominent by 35 msec. By 40 msec a large part of the epicardial surface of the left ventricle has been activated as has much of the right ventricular free wall creating a larger electrical hole in the wave front manifested on the surface by the prominent anterior sink.^{2, 3}

At 20 msec in the obstructive group the anterior bulge of positivity is much more prominent and reveals a more significant rightward component reflecting the individuals in this group who had some degree of right ventricular outflow tract obstruction and right ventricular free wall enlargement as well as simply septal hypertrophy.^{2, 4}

By 30 msec in the nonobstructive group the prominent negative sink the surface representation of intercostal breakthrough on both sides of the septum is a prominent feature while in the obstructive group there is continuation of the prominent outward bulging mound of positivity which has now moved slightly more leftward. This mound is increased in amplitude and is prolonged in duration over the normal. It is in distinct contrast to the surface sink of negativity which is expected as left ventricular free wall activation proceeds laterally and posteriorly after the electrical front has opened due to epicardial breakthrough of the activation wave in the region of the anterior septum. The non-obstructive map in general follows the normal progression with greater prominence of the negative sink which is both deeper and wider. This reflects the increased size of the activation front at the time of epicardial breakthrough to the surfaces of the enlarged and more widely separated right and left ventricles.

Individual maps of a 41 year old man with nonobstructive cardiomyopathy and a 30 year old man with IHSS make the points much more clearly since averaging has not attenuated detail (Fig. 2). At 10 msec a slight bulge of positivity is seen

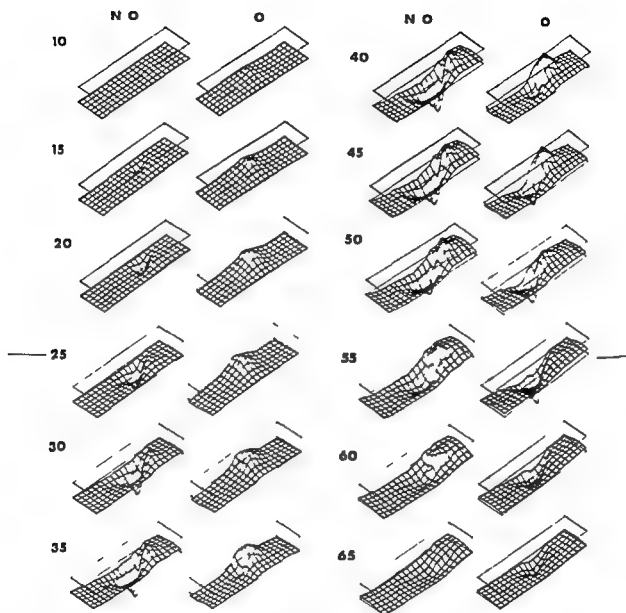


Fig 2 At intervals of 5 msec individual map from two patients with hemodynamically distinct form of cardiomyopathy are illustrated. Under the heading NO are seen the map of a 41 year old man with a history of regular alcohol intake on a daily basis for the last 21 years, left ventricular enlargement and failure with episodes of acute pulmonary edema and the continued evidence of left ventricular dysfunction in the form of S_1 and S_4 gallops, an elevated left ventricular end diastolic pressure and decreased indices of contractility. The patient whose map are seen under the heading O is a 30 year old man with IHSS with typical physical findings of a rapid rise of the carotid pulse stroke, a prominent bifid arterial pulse, an S_4 gallop, a systolic ejection murmur along the left sternal border without an ejection click which intensifies on Valsalva maneuver and with amyl nitrite and decreases with raising the leg. There were typical findings at cardiac catheterization including a subvalvular chamber with a gradient demonstrated from the low left ventricle. The early rise of positivity is more prominent at 10 msec in the obstructive patient. By 15 msec there is some negative expression as early septal and right free wall breakthrough is expressed in the nonobstructive patient. The mound of positivity in the patient map with IHSS increases in magnitude and moves to the right by 15 msec. Its further increase in positivity is contrasted with the increasing negativity and rather attenuated positivity in the nonobstructive patient at 20 msec. The positive mound increases and moves leftward through 30 msec with the earliest evidence of right ventricular breakthrough not occurring until 35 msec in the patient with IHSS. Negativity has been manifested in the nonobstructive patient to some extent since 15 msec and reaches its peak by 40 msec with not only prominent magnitude but a very large circumference. Left negativity in the obstructive patient is not reached until 55 msec, of lesser magnitude and has a less impressive perimeter. Pericardial positivity on the other hand occurs later in the nonobstructive patient and is more posteriorly oriented.

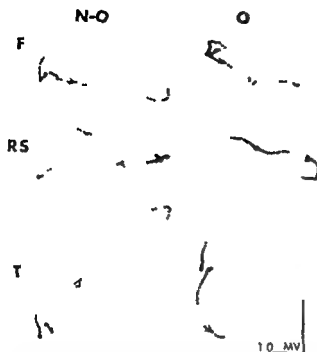


Fig. 3 The left column contains the frontal (F) right sagittal (RS) and transverse (T) loops of the 41 year old man with nonobstructive cardiomyopathy while the right hand column demonstrates the comparable loop in the 30 year old man with HISS. Note the equatorial but posterior orientation of the loops in the patient with nonobstructive disease which is in distinct contrast to the relatively normal frontal plane axis well below the horizontal demonstrated in the patient with HISS. The large rounded transverse loop of the patient with HISS with a significant portion of the loop anterior as well as posterior is in contrast to the posterior figure of 8 in the transverse plane in the case of nonobstructive disease. The dots are at 1 msec intervals and the arrows indicate the direction of the loop at that point. Note the lack of normal early left to right activation in the patient with nonobstructive disease; the early loop slowing, noted in all planes in that patient as well as in the patient with obstructive disease. In each instance the arrows are located beyond 24 msec as determined programmatically from 112 sampling sites from the originally stored digital data. Thus two forms of early activation abnormalities are demonstrated: (1) the failure of normal dominant left to right activation seen in the patient with nonobstructive cardiomyopathy and (2) an abnormal degree of early loop slowing, best demonstrated in the patient with HISS. There was no full *c* infarction pattern present in these instances.

in the nonobstructive patient. This bulge is more prominent in the obstructive group and has moved more rightward reflecting the cumulative activation of the septum and the right ventricle near its septal attachment. By 15 msec in the nonobstructive patient some surface negativity is already apparent as perhaps early right

ventricular breakthrough has occurred and has opened the front while, in contrast, in the obstructive patient the dominant positive pattern remains and continues to increase. By 25 msec the anterior opening in the electrical front is becoming more prominent on the surface at the time the lateral left ventricular muscle mass is being activated in the nonobstructive patient while positivity continues to summate in the obstructive patient. It is not until 35 msec in the patient with HISS that a very attenuated sink of negativity appears suggesting that septal and right ventricular breakthrough have occurred. The perimeter of negativity as well as its maximum magnitude is contrasted in the two patients at 40 and 45 msec while at 30 and 35 msec the maximum negative dip in the obstructive patient occurs.

In Fig. 3 VCG loops and scalar leads of the same two individuals are illustrated. The equatorial orientation of the VCG loops present in the patient with nonobstructive disease was characteristic of the entire group with the mean axis hovering close to the transverse plane with very little deviation along the Y axis. In contrast a much larger percentage of the vector loops in the obstructive group had their mean spatial axes slightly below the horizontal while the transverse loop was almost always large and round with a prominent anterior as well as posterior contribution as seen in the patient illustrated.

Varying degrees of abnormality in the early part of ventricular activation were demonstrated in loops, scalar leads and body surface maps. The most subtle form of abnormality was frequently seen in the obstructive patients where there was abnormal slowing of the VCG loops in all planes through at least 24 msec. A slightly more apparent abnormality in early activation was seen in over 40 per cent of the nonobstructive patients in whom there was absence of evidence of early left to right septal activation in the scalar leads, VCG loops or body surface maps. This was manifested as an absent septal Q wave in the λ lead or in all of ECC Leads I, aVL, V₁ or V₆ or an attenuated or absent early anterior bulge of positivity in the body surface map. This could also be seen as a lack of the rightward compo-

ment of the early direction of the VCG loop. The most blatant form of abnormal early activation was seen in one third of our patients with obstructive disease in the form of abnormally prolonged Q waves usually found in the mid and lateral precordial leads but sometimes in other lead sets. Abnormality was defined as Q wave duration greater than 0.03 sec though the incidence of abnormality was unchanged when criteria of 0.04 sec or greater were applied.¹¹ Abnormal valleys of negativity accompanied these findings in the body surface map.

Discussion

As to the genesis of the faulty left to right activation across the septum necessary to explain certain of the abnormal findings in the early part of ventricular activation we still lack electrophysiologic detail. Septal fibrosis has been offered to explain the lack of normal septal Q but this has been difficult to substantiate morphologically except in about one third of the patients.¹²

As to the genesis of the delayed activation suggested in the obstructive group and the presence of frankly abnormal Q waves seen in certain obstructive patients the common explanation has been to relate the delayed breakthrough to the right side of the septum to septal hypertrophy. Whether this is the case or not is unclear but data obtained by van Dam and associates¹³ and Durrer³ in plunge electrode studies of patients with hypertrophic subaortic stenosis strongly suggest another explanation. This work suggests almost synchronous activation of over half of the thickness of the left ventricular free wall which is in contrast to the expected outward spread of the wave of activation seen in successive layers from endocardium to epicardium causing an electrical gradient with rapid building of the positive expression. The epicardial electrode in Durrer's and van Dam's cases continued to see cavity potential until a much later than expected time in the QRS complex while the plunge electrodes recorded the fact that near to the epicardial surface after simultaneous activation of the underlying subendocardial layers had occurred an activation gradient could be seen to begin then to falter and finally

to spread to the epicardium. If this general pattern of altered activation occurs with progressively increasing degrees of impairment in cardiomyopathy and indeed if it obtains in the interventricular septum as well as in the left ventricular free wall it might well explain not only the presence of abnormal Q waves but certain other manifestations of early abnormality of activation such as the early slowing of the loop seen in certain patients as well as the false infarction pattern seen in others. At this writing however there is no direct electrophysiologic data to support the occurrence of actual slowing of conduction or the presence of septal activation abnormality.

Summary

Extensive noninvasive electrophysiologic studies have been performed on patients with cardiomyopathy. Body surface potential maps in the nonobstructive patients mimicked the normal in the general progression of the surface manifestations of activation but demonstrated relatively more prominent patterns of electrical breakthrough in terms of absolute magnitude and in terms of the perimeter of negativity at a given instant in time. The pattern of the obstructive group was distinctly different being marked by increased magnitude and duration of positivity with an obvious delay in septal right ventricular and left ventricular breakthrough contiguous to the septum with attenuated negativity at the time of those breakthroughs. Maximum negativity also usually occurred 15 to 20 msec later in the patients with IHSS. The patients with nonobstructive disease characteristically had posteriorly and equatorially oriented VCG loops and often lacked evidence of normal early left to right activation of the interventricular septum. Loops of the patients with obstructive disease were usually oriented below the horizontal and were less posteriorly oriented. Additional abnormalities noted in some patients with obstructive disease included slowing of the loop through at least 24 msec. The most extreme form of abnormal sequence of ventricular activation was seen in one third of the patients with IHSS in the form of false infarction patterns. An electrophysiologic rationale of this latter abnormality has been sug-

gested by van Dijn and Durrer and is thought by us to represent the best current explanation.

We appreciate the referral of certain patients in this report by Drs. Martin J. Frank and Sankaran Vasokan as well as our Cardiology Fellows.

REFERENCES

- 1 Melice R and Pirunko A. An orthogonal lead system for clinical electrocardiography. *Am Heart J* 62:93 1961
- 2 Durrer D. Electrical aspects of human cardiac activity: a clinical physiological approach to excitation and stimulation. *Cardiovasc Res* 2:1 1968
- 3 Taccardi B. Distribution of heart potentials on the thoracic surface of normal human subjects. *Circ Res* 12:341 1963
- 4 Liddlemon C O, Ruester V J, Horan I G and Brody D A. Distribution of heart potentials on the body surface in five normal young men. *Am J Cardiol* 21:860 1968
- 5 Horan I G, Flowers N C and Brody D A. Principal factor waveforms of the thoracic QRS complex. *Circ Res* 15:131 1964
- 6 Pearson R B, Gillespie I I and Selvester R H. On line digital collection and display of total body surface ECG data. in Hoffman I, Hamby R and Glassman E. Editors. *Vectorcardiography*, ed 2. Amsterdam 1971. North Holland Publishing Company, pp 146-153
- 7 Taccardi B, Musso E and de Ambroggi L. Potential fields of normal and ischemic hearts during rest, ventricular excitation and recovery. in Hoffman I, Hamby R and Glassman E. Editors. *Vectorcardiography*, ed 2. Amsterdam 1971. North Holland Publishing Company, pp 142-145
- 8 Spach M S and Barr R C. Physiological correlates and clinical application of isopotential surface maps. in Hoffman I, Hamby R and Glassman E. Editors. *Vectorcardiography*, ed 2. Amsterdam 1971. North Holland Publishing Company, pp 131-141
- 9 Spach M S, Bonnera J P, Barr R C, Wallace A M, Long L C, Gallie T M and Muney J M. Isopotential surface maps in children with varying types of right ventricular hypertrophy. in Proc VII Ann Colloquium on Vectorcardiography, Bratislava, Czechoslovakia 1966
- 10 Bonnera J P, Hill J D, Spach M S and Moore L N. The origin of surface potentials in right ventricular hypertrophy. in Proc VII Ann Colloquium on Vectorcardiography, Bratislava, Czechoslovakia 1966
- 11 Horan I G, Flowers N C and Johnson J C. The significance of the diagnostic Q wave of myocardial infarction. *Circulation* 43:478 1971
- 12 van Dijn R T, Roos J P and Durrer D. Electrical activation of ventricles and interventricular septum in hypertrophic obstructive cardiomyopathy. *Br Heart J* 31:100 1971

The effect of heart rate, acetylcholine, and vagal stimulation on antegrade and retrograde His-Purkinje conduction in the intact heart

P Jacob Varghese MD
Anthony N Damato MD
Sun H Lau MD
Masood Akhtar MD
Gustavus A Bobb
Staten Island N Y

Various anatomical studies have demonstrated the presence of cholinergic fibers in the ventricles.^{1,2} Acetylcholinesterase activity has also been demonstrated in the canine ventricle.³ However, the role of these fibers in the electrophysiology of the normal ventricle is not well understood. There have been conflicting reports in the literature. Earlier workers have shown that both vagal stimulation and small doses of acetylcholine have no effect on the transmembrane action potentials of the Purkinje fibers and the ventricular myocardium.^{4,5} Later reports have suggested that both vagus and acetylcholine may have an effect on the automaticity, conduction and morphology of the action potentials of the specialized conduction tissue below the AV junction.⁶⁻¹¹ The present study was undertaken to evaluate the effect of acetylcholine as well as graded vagal stimulation and heart rate on both antegrade and retrograde His-Purkinje conduction.

Methods

Ten adult mongrel dogs (15 to 35 kilograms) were anesthetized with intravenous pentobarbital sodium (30 mg per kilogram of body weight). The animals were intubated and artificially ventilated by a Harvard respirator. The heart was exposed through a right lateral thoracotomy at the fourth right intercostal space. As previously described, close bipolar plunge electrodes were used to record electrograms from the region of sinus node (SN), Bachmann's bundle (BB), the right atrial appendage (RAA), the left atrial appendage (LAA), the posterior portion of the left atrium (LAI) and the coronary sinus (CS). Two sets of wires were inserted into the region of the bundle of His.¹² One set was used to stimulate and the other pair was used to record the electrical activity of the bundle of His. Additional wires were inserted into the right and left ventricles for pacing. The right cervical vagus was isolated and sectioned. A set of plunge wires

From the Department of Cardiology, St. Vincent's Hospital, Staten Island, N.Y.
Received for publication July 15, 1972.
Revised manuscript accepted for publication September 15, 1972.
Address reprint requests to Dr. Varghese, MD, Department of Cardiology, St. Vincent's Hospital, 111th St., Staten Island, N.Y. 10314.

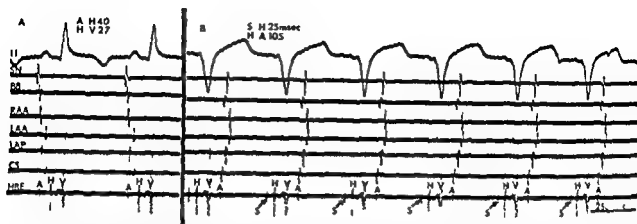


Fig 1 Panel A shows two sinus beats. In panel B the ventricle was paced at a cycle length of 330 msec. Note the 1:1 retrograde conduction to the atrium. Retrograde His-Purkinje conduction (S-H) measured 75 msec. SN = sinus node, BB = Bachmann's bundle, RAA = right atrial appendage, LAA = left atrial appendage, LAP = left atrium posterior, CS = coronary sinus, HBE = His bundle electrogram, I = low atrial septal electrogram, H = His deflection, V = ventricular electrogram, S = stimulus artifact. The above abbreviations are followed in Figs 2 to 9.

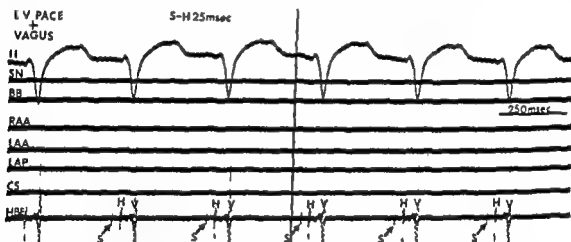


Fig 2 The ventricle was paced at the same cycle length as in Fig 1. Note the absence of atrial activity as a result of vagal stimulation. Retrograde His-Purkinje conduction time remained the same as in Fig 1.

was inserted into the distal stump in order to stimulate the vagus nerve. A battery-powered pacemaker* which delivered impulses of 2 msec duration at approximately $1\frac{1}{2}$ to 2 times threshold was used to stimulate the bundle of His or ventricles. The vagus nerve was stimulated by a Grass stimulator (Model S 8). The intensity of vagal stimulation was adjusted to produce a graded response varying from slowing of the sinus rate to cardiac arrest with and without atrial fibrillation. This was accomplished by delivering 2 msec impulses at a frequency of 40 impulses per second and at a varying voltage of up to 15 volts. In each study the following procedures

were performed. Both the His bundle and the ventricle were paced at varying cycle lengths ranging from 400 to 240 msec. The tracings were repeated with vagal stimulation and acetylcholine injection. Acetylcholine was injected into the femoral vein at a rate of 0.05 mg per kilogram per minute and this resulted in slowing of the sinus rate, increase in the A-V nodal conduction time followed by atrial asystole. All records were taken on a multichannel oscilloscopic photographic recorder at a paper speed of 200 mm per second.*

The following intervals were measured and expressed in milliseconds (msec). The A-H interval was measured from the onset

*Medtronic Inc (Model 5837) Minneapolis, Minn.

*Electronics for Medicine, White Plains, N.Y.

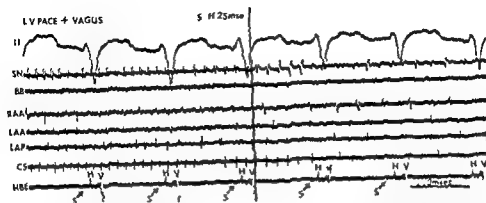


Fig. 3 Vagal stimulation resulted in atrial fibrillation and complete heart block. Ventricular pacing at the same cycle length as in Figs. 1 and 2 demonstrated similar retrograde His-Purkinje conduction time ($S-H = 15$ msec)

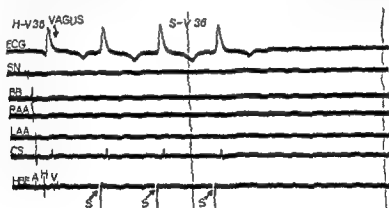


Fig. 4 The lack of effect of vagal stimulation on antegrade His-Purkinje conduction (see text)

of the low atrial depolarization (A) to the onset of the His deflection (H). The S-A interval during bundle of His pacing was measured from the stimulus artifact to the low atrial septal electrogram of the retrograde atrial impulse (A). The H-A interval during ventricular pacing was measured from the retrograde His deflection to the low atrial septal electrogram of the retrograde atrial impulse. Both the S-A and H-A intervals were considered to represent retrograde conduction time through the A-V node.

Antegrade His-Purkinje conduction time was measured from the His deflection to the onset of ventricular activation during sinus rhythm (H-V) or from the stimulus artifact to the onset of ventricular activation during bundle of His stimulation (S-V). During ventricular pacing the interval from the onset of the stimulus arti-

fact to the onset of the retrograde His deflection was used as a measure of retrograde His-Purkinje conduction time (S-H). Validation of the retrograde His deflection was obtained by demonstrating that the H-A interval during ventricular pacing was the same as the S-A interval during His pacing.

Results

Vagal stimulation. In all ten dogs the effects of vagal stimulation on antegrade and retrograde His-Purkinje conduction were evaluated during bundle of His and ventricular pacing at cycle lengths (CL) varying from 400 to 240 msec. At all CL of pacing vagal stimulation had no effect on either antegrade or retrograde conduction as compared to control. Representative examples are presented in Figs. 1 to 3 (same experiment).

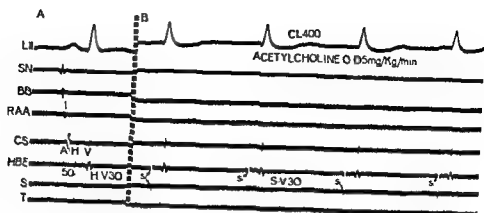


Fig. 5 The effect of acetylcholine on antegrade His-Purkinje conduction time

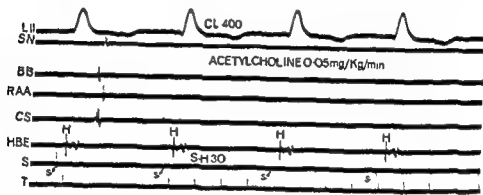


Fig. 6 Acetylcholine in doses sufficient to produce cardiac arrest had no effect on retrograde His-Purkinje conduction

Panel A of Fig. 1 illustrates two sinus beats in which the A-H and H-V intervals are 40 and 27 msec, respectively. In panel B the left ventricle was stimulated at a CL of 330 msec. One to one retrograde conduction to the atria occurred. Retrograde His-Purkinje conduction time (the interval between the stimulus artifact and the retrogradely activated His bundle deflection or S-H interval) measured 25 msec. Retrograde A-V nodal conduction time as indicated by the H-A interval was 10 msec.

In Fig. 2 vagal stimulation sufficient to cause cardiac arrest was applied. The left ventricle was stimulated at the same CL as in Fig. 1. Note the absence of atrial activity. Retrograde His-Purkinje conduction time (S-H interval) remained at 25 msec.

Vagal stimulation first resulted in cardiac arrest followed by atrial fibrillation with complete A-V block (Fig. 3). The left ventricle was then stimulated at a cycle length of 330 msec. Retrograde His-Purkinje conduction time remains constant at 25 msec.

The effects of vagal stimulation on ante-

grade His-Purkinje conduction are illustrated in Fig. 4. The first beat is the last of a series of sinus beats. The interval from the antegrade His deflection to the onset of the ventricular depolarization (H-V interval) measured 36 msec. Vagal stimulation resulted in cardiac arrest as evidenced by the absence of spontaneous atrial activity. Immediately after vagal stimulation was applied the His bundle was stimulated for three beats at a CL of 315 msec. The S-V interval was the same as the H-V interval during sinus rhythm. Following cessation of His bundle pacing the continued effects of vagal stimulation are noted.

Effect of acetylcholine The effects of acetylcholine infusion on antegrade and retrograde His-Purkinje conduction were also evaluated during bundle of His and ventricular pacing at cycle lengths varying from 400 to 240 msec. At all cycle lengths of pacing acetylcholine had no effect on either antegrade or retrograde His-Purkinje conduction as compared to control. Representative examples are presented in Figs. 5 and 6 (same experiment).

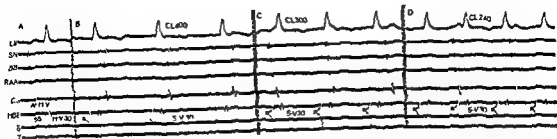


Fig. 7 Antegrade His-Purkinje conduction is not influenced by pacing the bundle of His at cycle lengths varying from 400 to 240 msec (see text)

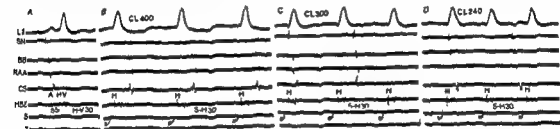


Fig. 8 Panel 4 shows a single sinus beat with A-H and H-V intervals of 55 and 30 msec respectively. During ventricular pacing (panels B to D) at decreasing cycle lengths the S-H interval remains constant at 30 msec

Fig. 5 shows the effect of acetylcholine on antegrade His Purkinje conduction time. A sinus beat with an H-V interval of 30 msec is shown in panel A. In panel B cardiac arrest occurred following injection of acetylcholine and bundle of His pacing at a cycle length of 400 msec resulted in an S-V interval which was the same as the normal H-V interval of the sinus beat (30 msec).

In Fig. 6 the right ventricle was paced at a cycle length of 400 msec. The first beat shows retrograde activation of the atria. Following this atrial arrest occurred as a result of acetylcholine injection. Retrograde His Purkinje conduction time (S-H 30 msec) was the same before and after acetylcholine.

Effect of heart rate. In all animals both the His bundle and the ventricle were paced at cycle lengths varying from 400 to 240 msec to evaluate the influence of the heart rate on the antegrade and the retrograde His Purkinje conduction. In addition retrograde His Purkinje conduction was also evaluated at cycle lengths as short as 160 msec.

Panel 1 of Fig. 7 demonstrates that normal antegrade His Purkinje conduction time (H-V) measured 30 msec. Panels B to

D show that antegrade His Purkinje conduction time (S-V 30 msec) remained constant during bundle of His pacing at cycle lengths varying from 400 to 240 msec. Fig. 8 demonstrates that ventricular pacing between cycle lengths of 400 to 240 msec had no effect on retrograde His Purkinje conduction time.

However when the CL of ventricular pacing was decreased to 200 msec or less retrograde His Purkinje conduction delay and block were noted as shown in Fig. 9. Panel A shows a control sinus beat with an H-V interval of 35 msec. In panel B the ventricle was paced at a cycle length of 200 msec. There was a 1:1 retrograde conduction to the bundle of His with a constant S-H interval. In panel C the cycle length of pacing was reduced to 190 msec. This resulted in 2:1 retrograde block to the bundle of His alternating with Wenckebach type of block. In panel D a fixed 4:1 retrograde block to the bundle of His occurred at a paced ventricular cycle length of 160 msec.

Discussion

The present study demonstrated that both vagal stimulation and acetylcholine have no effect on antegrade or retrograde

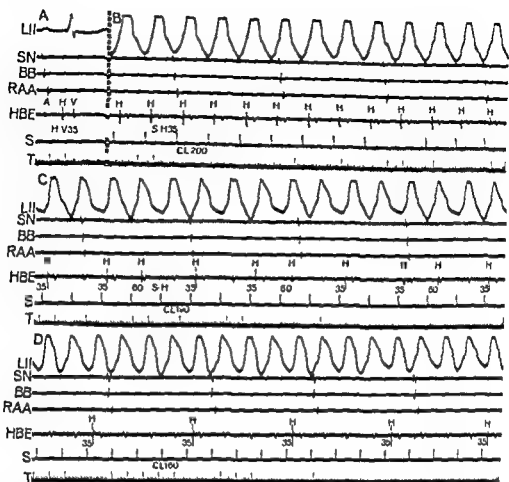


Fig 9 Panel A shows a sinus beat with integrated His-Purkinje conduction time of 35 msec. Panel B to D show the effect of ventricular pacing, at cycle lengths varying from 200 to 160 msec, on retrograde His bundle conduction (see text).

His-Purkinje conduction time. This is in marked contrast to their profound effects on A-V nodal conduction.¹⁸⁻²⁰ Our results support the findings of Alanis and associates²¹ who demonstrated that antegrade His-Purkinje conduction (H-V interval) was unaffected during vagal induced cardiac slowing. The present findings are also consistent with the observations that vagal stimulation and acetylcholine have no effect on the transmembrane action potentials of Purkinje and ventricular muscle fibers.²² Recently Bailey and associates²³ showed that superfusion with acetylcholine enhanced conduction in spontaneously firing Purkinje fibers. They demonstrated that with the infusion of acetylcholine the cells became less automatic, take off potential increased, rate of rise of phase 0 was enhanced, and conduction became more rapid. However, this enhancement of conduction with acetylcholine was shown in fibers with spontaneous phase 4 depolarization.

The results of the present study are also in agreement with the clinical observation that the electrocardiogram during effective carotid sinus massage, apart from slowing the atrial rate and producing some A-V nodal delay, fails to show any change in the duration and morphology of the QRS complex. The lack of QRS changes implies that antegrade His-Purkinje conduction is unaffected. A remote possibility is that His-Purkinje conduction is uniformly affected and consequently a change in the QRS complex would not be evident. This appears unlikely in view of the evidence of the present study.

In both the experimental animal and man progressive increases in the rate of atrial stimulation result in a progressive prolongation of A-V nodal conduction time.^{9,17,18} Similarly a progressive increase in retrograde A-V nodal conduction time occurs as the rate of His bundle or ventricular stimulation is increased.¹ In this respect the His-Purkinje system^{24,25} dif-

fers from the A V node. Both antegrade and retrograde His Purkinje conduction times are not affected by increasing rates of atrial or ventricular stimulation. This lack of effect is due to the influence of heart rate on the refractory periods of the HPS and the myocardium.^{21, 22} With increasing rates the refractory periods of both the HPS and ventricular myocardium are progressively shortened and as a result His Purkinje conduction remains unaffected but this response is true only up to certain heart rates. With very fast rates (cycle length 190 to 160 msec) various degrees of retrograde block to the HPS occurred. The site of block may be at the Purkinje muscle junction or at any site along the HPS wherever the site of maximum refractoriness occurs.

The mechanism of this type of block between the myocardium and the HPS can be explained as follows. At very fast rates instead of progressive shortening of the refractory period of both the HPS and the myocardium a dissociation of the refractoriness of these tissues may occur. While the refractory period of the myocardium shortens the refractory period of the HPS may remain the same. This would result in various types of block and may set up a situation for re entry.

The observation of retrograde block into the HPS at very fast ventricular stimulation may have some clinical significance. Since dissociation of the refractoriness between the myocardium and the HPS occurs at very fast rates one might speculate that the same phenomenon may occur with a very closely coupled ventricular premature beat. If this premature beat is sufficiently delayed in the HPS then some areas of the ventricular myocardium might be recovered and be ready to be depolarized again. Thus an intraventricular re entry could occur which might give another explanation for some forms of ventricular tachycardia and ventricular fibrillation initiated by a closely coupled ventricular premature beat.

Summary

In ten dogs the effects of acetylcholine graded vagal stimulation and heart rate on both antegrade and retrograde His Purkinje conduction were evaluated. His bundle pacing was used to obtain the antegrade His Purkinje conduction time and

pacing the ventricle and recording the retrograde His bundle electrogram gave the retrograde His Purkinje conduction time. Acetylcholine and vagal stimulation sufficient to produce cardiac arrest had no effect on antegrade and retrograde His Purkinje conduction. Similarly pacing at cycle lengths varying from 400 to 200 msec had no effect on antegrade or retrograde conduction time. However when the cycle length of ventricular pacing was decreased to less than 200 msec various degrees of retrograde His Purkinje conduction delay and block developed.

REFERENCES

1. Noorder J F. Studies on the innervation of the heart. I. Distribution of the cardiac nerves with special reference to the identification of the sympathetic or parasympathetic post ganglionic. *Am J Anat* 62:361 1939.
2. Stottler W A and McMahon R A. Innervation and structure of the conductive system of the human heart. *J Comp Neurol* 87:156 1947.
3. Teheng H T. Innervation of the dog's heart. *Am Heart J* 41:312 1951.
4. Mitchell J A G, Brown R and Cookson F B. Ventricular nerve cells in mammals. *Nature* 172:812 1953.
5. Napolitano J M, William V L, Hanlow C R and Cooper T. Intrinsic innervation of the heart. *Am J Physiol* 208:455 1965.
6. Cooper T. Terminal innervation of the heart. In Randall W, editor. *Nervous control of the heart*. Baltimore 1965. The Williams and Wilkins Company.
7. Jacobowitz D, Cooper T and Barner H. Histochemical and chemical studies of the localization of adrenergic and cholinergic nerves in normal and denervated cat hearts. *Fed Proc* 25:383 1966.
8. Brooks C, McC Hoffman M F, Suckling E E and Ona O. Excitability of the heart. New York 1955. Grune & Stratton Inc.
9. Hoffman M F and Cranefield P F. *Electrophysiology of the heart*. New York 1960. McGraw Hill Book Company Inc.
10. Eliaz M M, Bellet S, Tawil M and Muller O. Effects of vagal stimulation and acetylcholine on the ventricle. Studies in dogs with complete atrioventricular block. *Circ Res* 9:1372 1961.
11. Bailey J C, Greenspan H, Eliaz M V, Anderson G J and Fisch C. Effects of acetylcholine on automaticity and conduction within the canine proximal His Purkinje specialized conducting system. *Circ Res* 30:210 1972.
12. Hoffman M and Suckling E M. Cardiac cellular potentials. Effect of vagal stimulation and acetylcholine. *Am J Physiol* 173:312 1953.
13. Sano T, Ida Y and Hiraoka M. Action of acetylcholine on the Purkinje fiber studied

- by voltage clamp technique *Jap J Physiol* 20:155 1970
- 14 Scherlag B J, Kosowatz H D and Dimuto A N Technique for ventricular pacing from the His bundle of the intact heart *J Appl Physiol* 22:584 1967
 - 15 Dimuto A N, Lau S H and Bobb G A Studies on ventriculoatrial conduction and the reentry phenomenon *Circulation* 41:423 1970
 - 16 Lewis L The effect of vagal stimulation upon atrioventricular rhythm *Heart* 5:247 1914
 - 17 Alams J, Conzales H and Lopez E Electrical activity of the bundle of His *J Physiol* 112:127 1958
 - 18 Sherf D and Cohen J The atrioventricular node and selected cardiac arrhythmias New York 1960 Grune & Stratton Inc p 12
 - 19 Moe G K, Preston J B and Burlington H Physiologic evidence for a dual AV transmission system *Circ Res* 1:357 1956
 - 20 Dimuto A N, Lau S H, Bobb G A and Wit A L Recording of the AV nodal activity in the intact dog heart *Am Heart J* 80:331 1970
 - 21 Alams J, Lopez E, Madok J J and Pilar C Propagation of impulses through the atrioventricular node *Am J Physiol* 19:111 1959
 - 22 Hoffman B F and Suckling E E Effect of heart rate on cardiac membrane potentials and the unipolar electrogram *Am J Physiol* 179:123 1954
 - 23 Moore L N, Preston J B and Moe G K Duration of transmembrane action potential and functional refractive periods of canine atrioventricular and ventricular myocardium Comparison in single fibers *Circ Res* 24:251 1969
 - 24 June M J, Van der Steen A B M, Van Dam R Th and Durrer D Refractory period of the dog's ventricular myocardium following sudden changes in frequency *Circ Res* 24:251 1969
 - 25 June M J Effect of changes in heart rate on the refractory period of the heart Amsterdam 1971 Mondell Off setdrukkerij

Drug failure in reducing pressor effect of isometric handgrip stress test in hypertension

Sofjan Lamid MD

Frederick W Wolff MD*

Washington D C

Lind and associates¹ and Donald and associates² showed that sustained muscular contractions produced a marked increase of mean arterial pressure heart rate and cardiac output

Recently it was revealed that isometric handgrip contractions (IHC's) of patients with coronary artery disease³ with high left ventricle end-diastolic pressure⁴ and with ischemic heart disease⁵ produced higher increase of blood pressure than normals This simple technique of hand gripping has been investigated as a tool to detect abnormal left ventricle function^{6,7} The mechanism of action of this cardiovascular response to sustained exercises is not fully understood The present investigation was performed in order to study (1) the effect of IHC on the blood pressure in groups of normotensive subjects and untreated hypertensive patients and (2) the effect of a number of antihypertensive drugs with different mode of action on the blood pressure changes produced by IHC in hypertensive patients

Methods and subjects

In the first experiments two groups of subjects were used In one group were 12

normotensive volunteers five were male and seven were female three were Caucasian and nine were Negro their ages ranged from 20 to 82 years (mean = 50.42 S.E.M. = 4.01) Of these volunteers eight were normotensive patients of the Eye Clinic who were not taking medications and four were healthy employees of the Washington Hospital Center In the other group were 14 hypertensive patients (persons whose diastolic pressure had been more than 90 mm Hg on more than one occasion and without obvious renal or endocrine disorder as its cause) The ages of these patients ranged from 21 to 77 years (mean = 51.93 S.E.M. = 4.97) five were male and nine were female three were Caucasian and 11 were Negro Electrocardiograms (ECG) were made on all patients before the beginning of the experiment Hypertensive patients with systolic pressure of more than 200 mm Hg diastolic pressure more than 130 mm Hg with previous myocardial infarction angina pectoris stroke accelerated hypertension or heart failure were excluded from the study All patients were off medications for at least one month before entering the study There was no significant difference

From the Division of Medicine, Georgetown University School of Medicine, Washington, D.C.
Received July 1, 1972

Reprint requests to Sofjan Lamid, MD, Washington Hospital Center, 1101 Eye St. N.W., Washington, D.C. 20010.
Presented at the 1972 Meeting of the American Heart Association, Rockville, Md.

Table I ECG's of untreated hypertensive patients before entering the study

| ECG | No of patients |
|---|----------------|
| Within normal limits | 9 |
| Left ventricular hypertrophy | 3 |
| Left anterior superior hemiblock | 1 |
| Occasional premature ventricular contractions | 1 |

between the ages of hypertensive and normotensive subjects race and sex distributions of both groups were comparable

Technique of handgripping Before the onset of the experiment each subject was instructed in the technique of IHC test. After ten minutes in the sitting position blood pressure of the left arm was measured by a research nurse and then the subject pulled the *Handdy-namometer** with the right hand and the load of 25 per cent of the maximal voluntary contractions (MVC) was maintained for three minutes. Blood pressure was recorded at rest and at the end of three minutes of IHC.

In subsequent experiments the effects of chronic administration of several anti hypertensive drugs on the pressure response to IHC were measured. The 14 hypertensive patients were treated in a cross over trial according to a randomized block design.⁶ The following drugs were used: hydrochlorothiazide (25 mg three times a day), guanethidine (50 mg a day), reserpine (0.25 mg twice a day) and hydralazine (25 mg three times a day). All drugs were given orally and each period of treatment lasted four weeks. All patients were ambulatory with a normal diet and their blood pressure was recorded every two weeks at the same time of the day and in the same environments. Two blood pressure measurements were made as a base line prior to drug therapy. At each clinic visit the blood pressure response to IHC was evaluated. The average of two blood pressure measurements at two week intervals before and after treatments with each drug was computed and analyzed statistically.

Table II Mean blood pressure at rest of untreated hypertensive patients (first visit) and normotensive subjects

| | Mean blood pressure at rest (mm Hg) | |
|--------------|-------------------------------------|------------|
| | Systolic | Diastolic |
| Hypertensive | 153.00 | 100.85 |
| SEM | ± 5.89 | ± 3.80 |
| Normotensive | 132.67 | 86.00 |
| SEM | ± 3.00 | ± 2.17 |
| t | 3.08 | 3.40 |
| P | <0.01 | <0.01 |

Statistical evaluation was performed in two ways for the first set of experiments where two groups were compared the *t* test was utilized. In the second set of experiments the effect of several drugs on IHC was measured by analysis of variance.

Results

Table I lists the ECG's of 14 untreated hypertensive patients before entering the study. The mean systolic and diastolic pressure at rest in hypertensive and normotensive subjects is described in Table II. There was a highly significant difference of blood pressure at rest between hypertensive and normotensive subjects. The percentage of blood pressure increase during IHC was calculated from the initial resting pressure. It is shown in Table III that the percentile increase of systolic and diastolic pressure after three minutes of exercises was significantly higher in the hypertensive group while the mean load of IHC differed nonsignificantly. Three hypertensive patients who had electrocardiographic evidence of left ventricular hypertrophy did not have higher response to IHC than the other eleven hypertensives.

In subsequent experiments we have shown that hydrochlorothiazide, guanethidine, reserpine and hydralazine at fixed dosage during four weeks reduced the systolic and diastolic pressure of hyperten-

*Smedley Handdy-namometer from Stoelting Company, Chicago, Ill.

Table III Mean percentage of blood pressure increase after three minutes of IHC and the mean load of IHC in untreated hypertensive patients (first visit) and normotensive subjects

| | Mean blood pressure increase after 3 min of IHC (% of initial resting pressure) | | Mean load of IHC (kg) |
|------------------------|--|-----------------|--------------------------|
| | Systolic | Diastolic | |
| Hypertensive S.E.M. | 18.79 ± 2.2 | 21.35 ± 4.51 | 8.4 ± 0.59 |
| Normotensive S.E.M. | 6.91 ± 1.87 | 8.61 ± 2.00 | 8.00 ± 0.62 |
| t | 4.08 | 2.58 | 0.47 |
| P | < 0.01 | < 0.05 | > 0.5 |

N.S. = not significant

Table IV Effect of drug therapy on resting blood pressure of hypertensive patients (average of two visits)

| | Mean resting blood pressure (mm Hg) | |
|---------------------|--|-----------|
| | Syst. | Diastolic |
| Before drug therapy | 154.71 | 101.93 |
| After drug therapy | | |
| Hydrochlorothiazide | 130.85 | 88.86 |
| Guaneethidine | 143.92 | 92.85 |
| Reserpine | 138.88 | 88.57 |
| Hydralazine | 135.85 | 91.15 |
| F | 3.32 | 3.40 |
| P | < 0.05 | < 0.05 |

sive patients significantly (Table IV). The antihypertensive effect of one drug differed nonsignificantly from the others. One patient could not tolerate guanethidine and he was considered a drug failure; one other patient was a drug failure on hydralazine.

Table V shows the effect of drug therapy on blood pressure increase to IHC in hypertensive patients and the mean load of IHC. There were no significant differences between the means of percentage increase of systolic and diastolic pressure after three minutes of IHC in patients prior to therapy and after treatments with hydrochlorothia-

zide, guanethidine, reserpine, and hydralazine. During these experiments the mean load of contraction (25 per cent MVC) was not significantly different.

Discussion

Many investigators have shown that psychological stress⁸ as well as cold pressor tests¹⁰ produced an abnormally large rise of blood pressure in hypertensive patients. Both the cold water stress and psychological stress tests depend upon altered vascular reactivity to the sensory inflow resulting from such stresses. The IHC on the other hand adds the dimension of work (intense muscular contraction).

In our test subjects hypertensive patients had a larger rise of blood pressure (systolic and diastolic) during IHC than did normotensive persons. This greater pressure response to IHC stress was true not only in absolute terms but also as percentage of the initial resting pressure. We agree with other investigators¹¹ that comparison of percentile changes of blood pressure is more desirable because it computes the initial stage of arteriolar reactivity. In our study most of the hypertensive patients had a normal ECG but in the absence of hemodynamic determinations we could not exclude any abnormal left ventricular function which would contribute to the presence of higher pressure response to IHC.

In the cross over trial in hypertensive

Table V Effect of drug therapy on the blood pressure increase after three minutes of IHC and the mean load during IHC in hypertensive patients (average of two visits)

| | Mean blood pressure increase after 3 min of IHC (% of initial resting pressure) | | Mean load of IHC (kg) |
|---------------------|--|-----------|--------------------------|
| | Systolic | Diastolic | |
| Before drug therapy | 16.54 | 16.33 | 8.56 |
| After drug therapy | | | |
| Hydrochlorothiazide | 13.55 | 14.60 | 7.85 |
| Guanethidine | 12.73 | 14.52 | 7.96 |
| Reserpine | 13.60 | 14.01 | 8.06 |
| Hydralazine | 12.63 | 10.84 | 7.80 |
| F | 0.63 | 0.58 | 0.34 |
| P | NS | NS | NS |

patients we have shown that hydrochlorothiazide, guanethidine, reserpine, and hydralazine reduced the resting blood pressure significantly but did not affect the pressure response to IHC. Shapiro¹² showed that therapeutic doses of chlorothiazide and reserpine failed to reduce the blood pressure response to cold pressor and psychological tests. He suggested that hyperreactivity of blood pressure response to several types of stimuli in hypertensives is a constitutional character of these patients. Using several antihypertensive drugs with different mechanisms of action, our findings give support to this hypothesis.

It is common clinical practice to measure blood pressure at rest preferably after several minutes in the supine position in a quiet room. There is already voluminous literature on the effect of varying stresses in the life of hypertensive patients which may well have an effect on their life expectancy and incidence of complications. The present study suggests the use of IHC as a standard test for investigation, follow-up, and management of hypertension.

The failure to reduce pressure response to IHC and other stress including psychological tests by presently available antihypertensive drugs, has important therapeutic implications. It underlines the need for a general review and advice on the life pattern of hypertensive patients as an adjunct to the therapeutic regimen. The Veterans Administration Cooperative Study

(Group on Antihypertensive Agents)¹³ recently reported the beneficial effect of therapy on the morbidity and mortality rates of even moderate to moderately severe hypertensive patients but could not ascertain the benefit with regard to the progression of coronary heart disease. Our present findings point to a limitation of drugs in the total management of hypertension and suggest further studies on pharmacologic means to deal with the hypertensive effect of stress.

Summary

A pilot study is reported on the isometric handgrip contraction (IHC) test comparing its effects on the blood pressure of normotensive and untreated hypertensive persons and evaluating the effect of antihypertensive drugs on this test. IHC caused a greater percentile increase of blood pressure in untreated hypertensive patients and this hyperactivity of systolic and diastolic pressure to IHC stress is not affected by therapeutic doses of hydrochlorothiazide, guanethidine, reserpine, and hydralazine.

REFERENCES

1. Lind A R, McNicol G W, and Donald K W. Circulatory adjustment to sustained (static) muscular activity. In Evans K and Anderson K, editors. Physical activity in health and disease. Baltimore, 1966. The Williams & Wilkins Company, p. 38.
2. Donald K W, Lind A R, McNicol G W,

- Humphreys P W Taylor H and Staunton H P Cardiovascular response to sustained (static) contraction *Circ Res* 20:1 1967
- 3 Houston J D Atkins J M and Blomqvist G Cardiovascular response to isometric forearm contraction *Clin Res* 18:170 1970
- 4 Helfant R H deVillia M A Allan M B and Meister S C The effect of isometric handgrip stress on left ventricular function *Clin Res* 19:320 1971
- 5 Jackson D H Isometric stress: a new testing method in ischemic disease *Clin Res* 19:321 1971
- 6 Atkins J M Matthews O A Houston J M Blomqvist G and Mullins C B Arrhythmias induced by isometric (handgrip) exercise and dynamic exercise *Clin Res* 19:303 1971
- 7 Spangler R O and Linguist V A A Comparison of isometric and dynamic exercise on left ventricular function in normal and diseased hearts *Clin Res* 11:120 1971
- 8 Woolf C M Principles of biometry Princeton 1968 M Van Nostrand Company Inc p 126
- 9 Lorimer A H Macfarlane P W Irovan C Duffy T and Lawrie T M V Blood pressure and catecholamine response to stress in normotensive and hypertensive subjects *Cardio-vasc Res* 5:169 1971
- 10 Shapiro A P Moutros S E and Knischer E Patterns of pressor response to noxious stimuli in normal hypertensive and diabetic subjects *J Clin Invest* 42:1890 1963
- 11 Gombos E A Hulet W H Bopp P Goldring W Baldwin D S and Chasis H Reactivity of renal and systemic circulations to vasoconstrictor agents in normotensive and hypertensive subjects *J Clin Invest* 41:203 1962
- 12 Shapiro A P Pressor response to noxious stimuli in hypertensive patients Effect of reserpine and chlorthalidate *Circulation* 26:242 1962
- 13 Veterans Administration Cooperative Study Group on Antihypertensive Agents Effects of treatment on morbidity in hypertension *JAMA* 213:1143 1970

The role of beta adrenoceptors in the coronary and systemic hemodynamic responses to emotional stress in conscious dogs

M Bergamaschi Ph D
A M Carataggi D I M
I Mandelli D Sc
R G Shanks M D *
Milan, Italy

Studies in man and in conscious dogs have shown that marked changes occur in the cardiovascular system during emotional stress.^{1,2} The response is characterized by increases in heart rate, arterial blood pressure and cardiac output and by a reduction in peripheral vascular resistance. The elegant studies by Rayford and associates³ also demonstrated marked changes in coronary hemodynamics in conscious dogs during emotional stress. There were increases in mean and stroke coronary blood flow and a marked reduction in coronary vascular resistance.

As the sympathetic nervous system plays an important role in mediating the cardiovascular responses to emotional stress, the development of drugs which specifically block beta adrenoceptors has provided a new and useful tool for analysis of the contribution of activity of these receptors to the hemodynamic reactions to stress. Propranolol has been shown to reduce the increases in heart rate and cardiac output and the decrease in peripheral vascular resistance during emotional stress in man.²

Oxprenolol almost completely abolished an emotional tachycardia in ski jumpers.⁴ Similar results have been obtained by Pitt and associates^{4,5} in conscious dogs undergoing stress. The changes in heart rate, arterial pressure, cardiac output, and peripheral vascular resistance were completely blocked by propranolol. In the same experiments propranolol did not alter the coronary hemodynamic response to stress.⁶ These results give further evidence that the systemic hemodynamic response to emotional stress is largely controlled by the sympathetic nervous system but do not clarify the role of beta adrenoceptors in mediating the changes in the coronary circulation occurring during emotional stress in conscious dogs.

As beta adrenoceptors blocking drugs have been reported to possess different degrees of action on beta receptors located in different tissues and additional properties—e.g. local anesthetic or quinidine like activity and intrinsic sympathomimetic activity^{7,8}—we decided to compare the effects of four beta adrenoceptor blocking

From the Carlo Erba Institute for Therapeutic Research, Milan, Italy.

Received for publication Nov. 28, 1972.

Reprint requests to Dr. Mario Bergamaschi, Institute Carlo Erba, Via Imbonati 24, 20159 Milano, Italy.

*Department of Therapeutics and Pharmacology, The Queen's University Belfast, Northern Ireland.

drugs propranolol sotalol (VIJ 1999) practolol (ICI 50172) and alprenolol (H 56/28) on the systemic and coronary hemodynamic responses to emotional stress in conscious dogs

Methods

Healthy mongrel dogs weighing 14 to 18 kilograms were used in these experiments. The surgical procedures and postoperative care of the animals were similar to those previously described¹⁰ and will only be described briefly. Anesthesia was induced by the intravenous injection of thiopentone 30 mg per kilogram of body weight and maintained by artificial respiration with oxygen to which halothane (2 per cent to 3 per cent) was added when required. The chest was opened through the fourth left intercostal space and the heart was exposed. Electromagnetic flow transducer (Biotronex Laboratories) were implanted around the left circumflex coronary artery and the ascending aorta. A pneumatic cuff was applied to the left coronary artery proximal to the flowprobe. Inflation of the cuff to occlude the artery was used to determine zero flow postoperatively. A polyethylene catheter filled with a solution of 0.9 per cent NaCl was inserted through a carotid artery until its tip was shown by radiocopy to be in the aortic arch.

Observations were not made until 15 days after insertion of the flow probes and aortic catheter when all dogs were in good health. Phasic aortic and coronary flow tracings were obtained by means of an electromagnetic flowmeter (Biotronex Laboratories). Phasic and mean aortic pressure were measured by a Statham P 23 Db transducer connected to the aortic catheter. Postoperatively the electrocardiogram was obtained by parasternal electrodes attached to the skin with adhesive tape. The electrocardiogram, aortic pressure and phasic and mean aortic and coronary flow were recorded on an 8 channel Beckman Type R Dynograph. Stroke volume and stroke coronary flow were measured by planimetric integration of the areas beneath the respective phasic flow patterns of at least 3 consecutive cardiac cycles.

The following parameters were calculated: mean systemic resistance (mean aortic pressure in mm Hg/cardiac output

in ml per minute) stroke resistance (mean arterial pressure in mm Hg/mean coronary flow in ml late diastolic coronary resistance was also calculated using the procedure of Rayford and colleagues⁸. Left ventricular work/time in kg m per minute was calculated as a product of cardiac output (liters per minute) and mean aortic pressure (mm Hg \times 13.6).

Emotional stress was induced in all dogs by an auditory stimulus. In order to standardize the stimulus a starting pistol the detonations of which were approximately of the same intensity on each occasion was used. The responses to excitement were determined four times in each dog on each day on two occasions 30 minutes apart during a control period and 5 minutes after the intravenous injection of 0.2 and 1.0 mg per kilogram of body weight of each drug. The results obtained from all the parameters were submitted to an analysis of variance which was carried out on the differences between the data recorded at zero time and those observed 2, 6 and 10 seconds after each pistol shot. A two way cross-classification was used: difference between groups (Control stress 1, Control stress 2 and 5 minutes after 0.2 and 1.0 mg per kilogram of body weight of each drug) and difference between dogs.

The following drugs were used: propranolol (Inderal, Imperial Chemical Industries Ltd), sotalol (VIJ 1999, Mead Johnson Ltd), practolol (ICI 50172, Eraldine, Imperial Chemical Industries Ltd) and alprenolol (H 56/28, Aptin, A B Hassle). All drugs were dissolved in a sterile solution of 0.9 per cent NaCl; the doses are expressed in terms of the salt. Dilutions of isoprenaline were made from the commercial stock solution.*

Results

Response to emotional stress The systemic and coronary hemodynamic responses to an auditory stimulus in a conscious dog are shown in Fig 1. After firing the gun the dog, although trained to lie on its left side, became restless and shivered. An increase in the rate of respiration was observed. Heart rate increased to a maximum of 250

*We are indebted to Dr S. J. H. G. To, St. ICI, for the gift of the drugs used in this study. Dr A. B. Hassle for alprenolol.

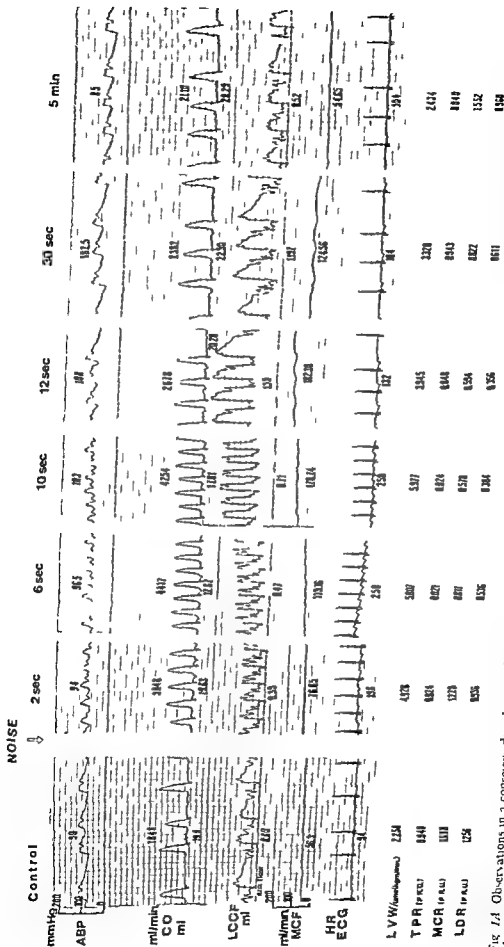


Fig. 1A Observations in a conscious dog, showing the effect of alprenolol on the response to stress. Records obtained before and at intervals after the firing of a pistol are: aortic pressure (ABP), phasic aortic flow (AO), phasic and mean left circumflex coronary flow (LCCF, MCF) and the electrocardiogram. The figures on the records, from above downwards, are mean arterial pressure, cardiac output, stroke volume, stroke coronary flow, mean coronary flow, and heart rate. The figures were obtained before the administration of alprenolol. Δ CR = total peripheral resistance, Δ VR = mean coronary artery resistance, LDR = left diastolic coronary resistance, LCOF = left coronary flow.

ALPRENOLOL (1 mg/kg) NOISE

5 min after 1 mg/kg LV ↓

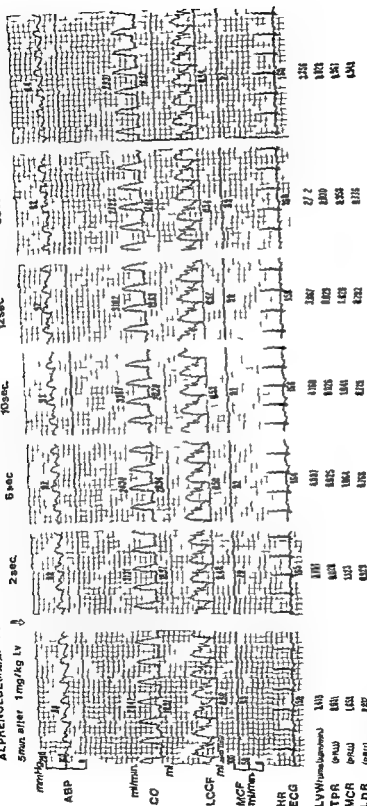


Fig. 12 Record of turn of a con cross dos. after the dos. of alprenolol 1 mg, per kilogram of body weight by intravenous injection for explanation see legend to Fig. 11

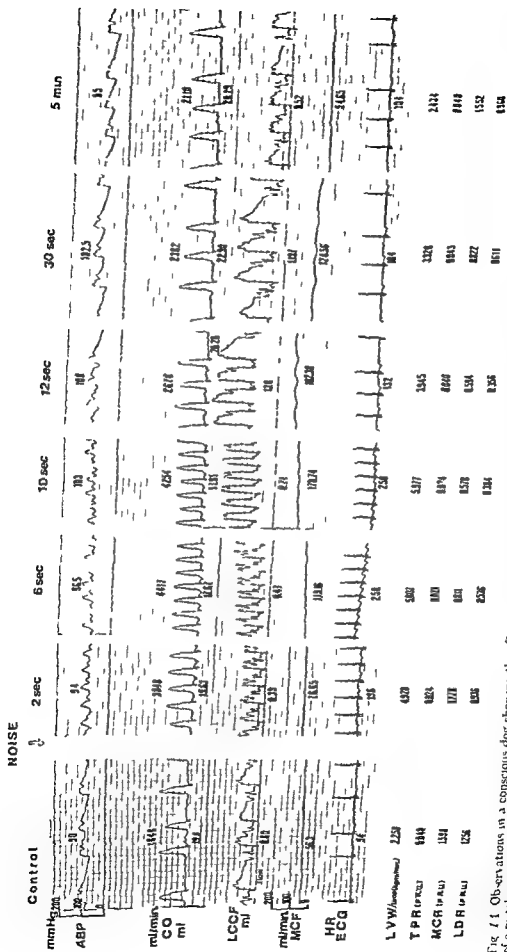


Fig. 11. Observations in a conscious dog showing the effect of alprenolol on the response to stress. Record obtained before and at intervals after the firing of a pistol are aortic pressure (ABP), phasic aortic flow (CO), phasic left circumflex coronary flow (LCCF), MCF, and the electrocardiogram. The figure shows the response to stress before and after the administration of alprenolol. The values are mean \pm SEM. The values in parentheses are the values obtained before the administration of alprenolol.

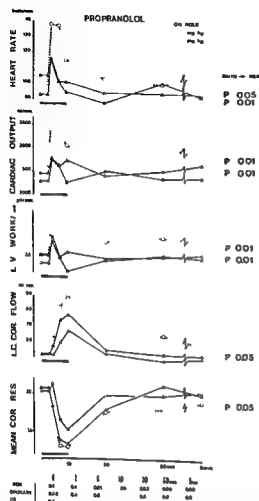


Fig 3 Effect of the intravenous injection of propranolol 0.2 and 1.0 mg per kilogram of body weight on the responses to stress in conscious dogs. Averaged results from 3 dogs at selected intervals before and after the application of stress on 4 occasions. Responses \circ — \circ mean of 2 sets of results before propranolol Δ — Δ result after propranolol 0.2 mg per kilogram of body weight \blacktriangle — \blacktriangle results after propranolol 1.0 mg per kilogram of body weight. Legend and determination of values for p were similar to those in Fig 2 with Δ and \blacktriangle referring to effects of propranolol 0.2 and 1.0 mg per kilogram of body weight respectively.

in mean and late diastolic coronary resistance and the increase in mean coronary flow produced by stress were significantly reduced after propranolol 0.2 mg per kilogram of body weight but they were unaltered after propranolol 1.0 mg per kilogram of body weight.

Effects of practolol The averaged results

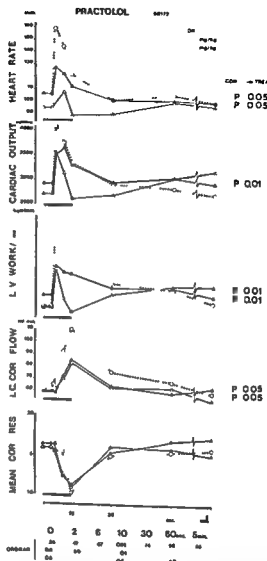


Fig 4 Effect of the intravenous injection of practolol 0.2 (Δ — Δ) and 1.0 (\blacktriangle — \blacktriangle) mg per kilogram of body weight on the responses to stress in conscious dogs. Averaged results from 3 dogs. Presentation of results and legends as in Figs 2 and 3.

from the three dogs which received practolol 0.2 and 1.0 mg per kilogram of body weight are given in Fig 4. Practolol 0.2 mg per kilogram of body weight significantly reduced the increases in heart rate, cardiac output, left ventricular work, and left coronary flow produced by stress. No further changes were produced by practolol 1.0 mg per kilogram of body weight. The increase in stroke coronary flow in response to stress before practolol was not

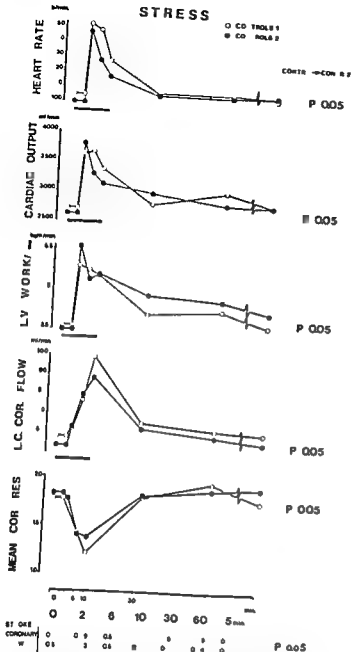


Fig 2 Responses to two stressful stimuli applied 30 minutes apart in 4 conscious dogs. The averaged results from the 4 dogs for heart rate, cardiac output, left ventricular work, left circumflex coronary flow, stroke coronary flow, and mean coronary resistance obtained at intervals after application of the stimulus are shown. Responses: \circ — \circ first stimulus; \bullet — \bullet second stimulus. Values for p were determined for the differences between the two runs using the accumulated results obtained during the period indicated by the solid bar (0, 2, 6, and 10 seconds).

beats per minute within 10 seconds and then declined rapidly. Mean aortic pressure increased during the same time from 90 to 108 mm Hg. Although stroke volume decreased, cardiac output was raised from 1,846 to 4,417 ml per minute due to the cardiac acceleration. As arterial pressure

and cardiac output were increased at the same time, the calculated external work of the left ventricle increased from 1.76 to 5.97 kg m per minute while total peripheral resistance decreased from 0.048 to 0.021 PRU.

Coronary flow was greatly increased in response to the stimulus and reached its highest value at 12 seconds which was 2 to 6 seconds after the peak values for heart rate and cardiac output. Systolic and diastolic coronary flow decreased 2 seconds after the stimulus and then increased so that stroke coronary flow increased from 0.60 to 1.38 ml at 12 seconds. The concomitant increase in heart rate contributed to an increase in mean coronary flow from 36 to 182 ml per minute when the mean coronary resistance fell from 1.59 to 0.59 PRU. Systolic, diastolic, and late diastolic coronary resistances were all reduced. The percentage of cardiac output entering the left circumflex coronary artery at rest was 3.05; it rose to a maximum of 6.81 twelve seconds after firing the gun. Five minutes after application of the stimulus all values except arterial pressure and cardiac output had returned to normal.

Similar changes were obtained in all dogs in response to the first stimulus on each day of observation. A second stimulus was applied about 30 minutes after the first when all values had returned to normal. There was no significant difference between the responses to these two stimuli (Fig 2).

Effects of propranolol. Observations were made in three dogs in which the responses to stress were elicited before (2 occasions) and after the intravenous injection of propranolol 0.2 and 10 mg per kilogram of body weight. The averaged results from the three dogs are given in Fig 3. The administration of propranolol (0.2 mg per kilogram of body weight) reduced heart rate, cardiac output, left ventricular work, and mean coronary flow. Stroke volume and stroke coronary flow were unaltered and mean and late diastolic resistance were increased. Propranolol 0.2 mg per kilogram of body weight significantly reduced the increases in heart rate, cardiac output, and left ventricular work which occurred in response to stress. No further change in the systemic responses occurred after propranolol 1 mg per kilogram of body weight. The reduction

and practolol produced small and not significant changes in resting systemic hemodynamics and in coronary blood flow and vascular resistance. This confirms earlier observations by Pitt and colleagues¹² and by Bergamaschi and co-workers¹³ which showed that in conscious dogs propranolol does not increase coronary vascular resistance. Alprenolol which had already been shown to possess intrinsic sympathetic activity¹⁰ significantly increased heart rate, cardiac output and left ventricular work but decreased systemic resistances.

Although the outward signs of emotional stress were still present after administration of all four drugs, the hemodynamic responses to stress were significantly altered. All drugs significantly reduced the increases in heart rate, cardiac output, aortic pressure and left ventricular work. These reductions were produced by the smaller dose (0.2 mg per kilogram of body weight) of propranolol, practolol and alprenolol but only by the largest dose (1 mg per kilogram of body weight) of sotalol.

It has been reported¹⁴ that the hemodynamic response to stress in the conscious dog is similar to that recorded during sympathetic stimulation. Subsequent experiments¹⁵ showed that the sympathetic nervous system is predominantly involved in the response to stress. The present results confirm that the systemic hemodynamic changes elicited by stress are largely mediated by beta-adrenoceptors as they were significantly altered by all four drugs. Similar results³ were obtained in pilots training in a flight simulator: the increases in arterial pressure, heart rate and cardiac output induced by this stressful situation were markedly inhibited by propranolol.

Although the hemodynamic responses to stress were significantly reduced after the administration of propranolol 1 mg per kilogram of body weight, changes still occurred in all parameters. The mechanism of these increases in heart rate, cardiac output and left ventricular work is not clear. They are unlikely to result from incomplete blockade of beta-adrenoceptors as several studies^{16, 17} have indicated that propranolol 1 mg per kilogram of body weight produces a high degree of blockade of these receptors. The increase in heart

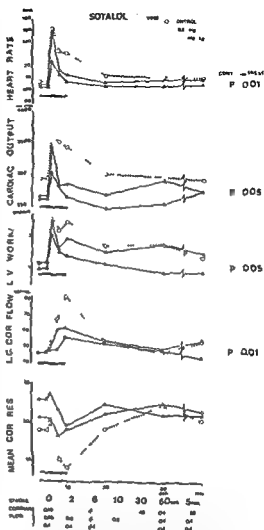


Fig 6 Effect of the intra-venous injection of sotalol 0.2 (Δ — Δ) and 1.0 (\blacktriangle — \blacktriangle) mg per kilogram of body weight on the response to stress in conscious dogs: averaged results from 3 dogs. Presentation of results and legends as in Figs 2 and 3.

rate in response to stress after propranolol may result from a reduction in vagal inhibitory activity to the heart, such a mechanism contributes to the increase in heart rate produced by exercise after the administration of propranolol¹⁸. However, it is not known whether the mechanism of the changes in cardiac function after propranolol in response to exercise and to emotional stress have a common origin. If they have, the increases in cardiac function produced by stress after beta-blockade may result from both withdrawal of activity of the

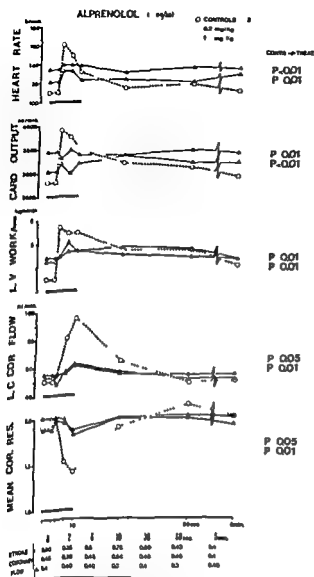


Fig 5 Effect of the intravenous injection of alprenolol 0.2 (Δ — Δ) and 1.0 (\blacktriangle — \blacktriangle) mg per kilogram of body weight on the responses to stress in conscious dogs. Averaged results from 3 dogs. Presentation of results and legends as in Figs 1 and 3.

altered after both doses of the drug. Similar reductions in mean and late diastolic resistance occurred in response to emotional stress before and after the two doses of practolol.

Effects of alprenolol The results of an experiment showing the response to stress before and after alprenolol 1 mg per kilogram of body weight, are shown in Fig 1. The responses before the drug have been described above. The administration of alprenolol increased heart rate, cardiac output, and left ventricular work. Although the increase in heart rate was accompanied by a decrease in stroke coronary flow, mean

flow through the left circumflex coronary artery was increased. Systemic and coronary resistance were slightly decreased. The hemodynamic responses to stress after the administration of alprenolol 1 mg per kilogram of body weight, were markedly altered. The increases in heart rate, cardiac output, left ventricular work, and coronary flow were greatly reduced. The reduction in systemic and coronary resistance were also inhibited. The averaged results from three dogs which received alprenolol 0.2 and 1 mg per kilogram of body weight, are given in Fig 5. The increases in heart rate, cardiac output, and left ventricular work produced by stress were significantly reduced after alprenolol 0.2 mg per kilogram of body weight and almost completely blocked after 1 mg per kilogram of body weight. The changes in coronary flow and resistance were reduced to the same extent.

Effects of sotalol The averaged results from the three dogs which were given sotalol, 0.2 and 1 mg per kilogram of body weight are shown in Fig 6. The administration of sotalol reduced heart rate, cardiac output, left ventricular work, mean and stroke coronary flow, with no change in stroke volume, mean and late diastolic coronary resistance were increased. The increase in heart rate, cardiac output and left ventricular work produced by stress were significantly reduced after sotalol 1.0 mg per kilogram of body weight but were largely unaltered by 0.2 mg per kilogram of body weight. The changes in mean and stroke coronary flow and in mean and late diastolic resistance in response to stress were also reduced by sotalol.

Discussion

The changes in systemic and coronary hemodynamics induced by stress in the present experiments are similar to those previously reported in conscious dogs.¹⁴ Delivery of an auditory stimulus significantly increased heart rate, cardiac output and left ventricular work, peak blood pressure rises were always accompanied by slowing of the heart rate, which is likely to depend upon baroreceptor reflex.¹¹ Stroke and mean coronary flow were increased and coronary resistances were decreased.

In the present experiments propranolol

propranolol also indicates that activation of beta adrenoceptors in the coronary vessels is involved in the physiological adaptation of the coronary circulation to a stressful situation

Summary

The present observations were made to investigate the role of beta adrenoceptors in the hemodynamic response to stress in conscious dogs. Probes for electromagnetic flowmeter were implanted on the ascending aorta and left circumflex coronary artery and a catheter was inserted into the aortic arch 15 days before observations were made. Emotional stress produced by firing a gun increased heart rate, cardiac output and left ventricular work. The responses were reduced by propranolol, practolol and sotalol 0.2 and 1.0 mg per kilogram of body weight and abolished by alprenolol. The difference between the effect of alprenolol and the other drugs may result from the marked intrinsic sympathomimetic activity of alprenolol which increased the resting level of heart rate, cardiac output and left ventricular work.

Emotional stress increased mean blood flow through the left circumflex coronary artery as a result of increase in stroke coronary flow and heart rate. Propranolol, sotalol and practolol reduced the increase in mean coronary flow produced by stress through a reduction in tachycardia; the increase in stroke coronary flow was unaltered. The decreases in mean and late diastolic coronary resistance produced by stress were not altered by these three drugs.

Alprenolol, unlike the other drugs, completely abolished the increases in mean and stroke circumflex coronary flow and the decreases in coronary vascular resistance produced by stress.

The present observations indicate that the dilatation of coronary arteries produced by stress does not result entirely from an increase in cardiac metabolism; they also suggest that stimulation of beta adrenoceptors in the coronary arteries plays an important role in the coronary vasodilatation produced by emotional stress.

The authors wish to express their gratitude to the staff members of the Cardiovascular Department of

the Carlo Erba Institute for their individual contributions to this study.

REFERENCES

- 1 Brod J. Haemodynamic basis of acute pressure reactions and hypertension. *Br Heart J* 25: 227 1963.
- 2 Elias H, Rosen A and Scott H M. Systemic circulatory response to stress of simulated flight and to physical exercise before and after propranolol blockade. *Br Heart J* 29: 671 1967.
- 3 Rayford C R, Khouri E M and Gregg D E. Effect of excitement on coronary and systemic energetics in unanaesthetized dogs. *Am J Physiol* 209: 680 1965.
- 4 Pitt M, Elliott E C, Khouri M and Gregg D E. Effect of adrenergic blockade on the coronary haemodynamic response to excitement at fixed ventricular rate in the unanaesthetized dog. *Physiologist* 9: 267 1966.
- 5 Imhof P R, Blatter K, Fuccella L M and Turri M. Beta blockade and emotional tachycardia: radiotelemetric investigations in ski jumpers. *J Appl Physiol* 27: 366 1969.
- 6 Pitt M. Sympathetic control of coronary circulation in the unanaesthetized dog. In Marchetti G and Taccardi B, editors. *Coronary circulation and energetics of the myocardium*. Basel 1967. S. Karger AG, pp 89-97.
- 7 Pitt M. Effect of propranolol on coronary haemodynamics in the unanaesthetized dog. In Hattiss A A, Ross G and Hall V E, editors. *Cardiovascular beta adrenergic responses*. Los Angeles 1970. University of California Press, pp 109-118.
- 8 Fitzgerald J D. Perspectives in adrenergic beta receptor blockade. *Clin Pharmacol Ther* 10: 292 1969.
- 9 Shanks R G. The properties of beta adrenergic blocking agents. *Irish J Med Sci* 2: 351 1969.
- 10 Bergamaschi M, Shanks R G, Caravaggi A M and Mandelli V. A comparison of the cardiovascular action of four adrenergic β receptor blocking agents in resting conscious dogs. *Am Heart J* 82: 338 1971.
- 11 Heymans C, Delaunoy A H and Van Den Heuvel Heymans G. Tension and distensibility of carotid sinus wall presensors and blood pressure regulation. *Circ Res* 1: 13 1953.
- 12 Pitt M, Greene H L, Sugishita Y and Ross R S. Effect of β -adrenergic receptor blockade on coronary haemodynamics in the supine unanaesthetized dog. *Cardiovasc Res* 4: 89 1970.
- 13 Granata L, Olsson R A, Huvos A and Gregg D E. Coronary inflow and oxygen usage following cardiac sympathetic nerve stimulation in unanaesthetized dogs. *Circ Res* 16: 111 1965.
- 14 Black J W, Duncan W A M and Shanks R G. Comparison of some properties of propranolol and propranolol. *Br J Pharmacol* 25: 577 1965.
- 15 Ledson J R, Linden R J and Norman J

vagus nerves and from an unknown mechanism.

In contrast to the effects of propranolol, alprenolol, 1 mg per kilogram of body weight, abolished almost completely the increases in heart rate, cardiac output, and left ventricular work produced by stress. As the beta adrenoceptor blocking activity of propranolol and alprenolol is similar¹⁰ the difference in their effects on the response to stress would appear to result from other differences in their properties. Alprenolol has intrinsic sympathomimetic activity¹⁰ which was apparent in the present experiments from the increases in resting levels of heart rate, cardiac output, and left ventricular work which it produced. As these increases were comparable to those produced by stress after propranolol, 1 mg per kilogram of body weight, they may have masked the effects of stress after alprenolol, 1 mg per kilogram of body weight, so that the inhibitory effects of the two drugs on the responses to stress may not be different. Both pinctolol and sotalol have been shown to be less potent than propranolol in blocking beta adrenoceptors. The results obtained in the present experiments confirm these previous findings as the two drugs were less effective than propranolol in antagonizing the systemic hemodynamic response to emotional stress.

Although propranolol reduced the increases in heart rate and cardiac work in response to stress, it did not significantly alter the intrinsic response of the coronary vessels to stress. Although the increase in mean and left circumflex coronary flow was reduced by the drug as a result of a decrease in the heart rate response, stroke coronary flow response to stress was unaltered. This conclusion is supported by the observation that the decreases in mean and late diastolic coronary resistances produced by emotional stress were not altered by propranolol. Previous studies⁷ have indicated that propranolol, although inhibiting the cardiac responses to stress in conscious dogs, did not alter the changes in the coronary circulation.

The effect of pinctolol was similar to that of propranolol. Sotalol significantly inhibited the increase in mean coronary flow and the reduction in mean and late diastolic

coronary resistance produced by stress.

Alprenolol almost completely blocked the coronary vascular response to stress as indicated by the lack of variation in stroke coronary flow and in mean and late diastolic coronary resistance after administration of the drug. The concurrent blockade of the cardiac acceleration prevented mean coronary flow from increasing. The reason for the difference between the effects of alprenolol and those of propranolol, pinctolol, and sotalol is not clear. It does not arise from differences in the beta blocking activity of alprenolol and propranolol since the two drugs were equally effective in blocking the effects of isoprenaline on the coronary vasculature in conscious dogs.¹⁰

The difference in the effects of the two drugs on coronary hemodynamics could result from alprenolol being more effective in blocking the effects of endogenously released noradrenaline although there are no other observations to support such a hypothesis.

Drugs which block beta adrenoceptors have been shown to inhibit the increases in cardiac work and oxygen consumption provoked by simulated exercise in anesthetized dogs¹⁷ and by exercise in conscious dogs¹⁸ but there are no reports of the effect of these drugs on the hemodynamic and metabolic changes induced by stress. In the present experiments propranolol and alprenolol (1 mg per kilogram of body weight) reduced the increases in work and rate of heart produced by stress by 71 and 75 per cent respectively. We did not measure the concurrent changes in oxygen consumption before and after beta blockade but as a relationship between cardiac work, heart rate, and oxygen consumption has already been demonstrated by previous authors^{12, 19} it can be assumed that in the present experiments the increases in oxygen requirement of the myocardium during stress were reduced to the same extent as the increase in cardiac work. The simultaneous observation that the coronary vasodilatation is unaltered by propranolol suggests that the changes induced by stress in the coronary circulation do not result entirely from an increase in myocardial metabolism. The blockade of the changes in coronary hemodynamics produced by al

Cardiac lymphangioma and lipoma

Report of a case of simultaneous occurrence in association with lipomatous infiltration of the myocardium and cardiac arrhythmias

Daniel T. Anbe M.D.
Gerald Fine M.D.
Detroit, Mich.

In spite of an increased awareness and improvement of diagnostic techniques primary tumors of the heart are infrequently encountered. The occurrence of multiple primary cardiac tumors is rarely reported and its incidence is unknown. While primary cardiac tumors may produce a variety of symptoms cardiac arrhythmias are relatively common in patients with metastatic or malignant primary tumors. They are less frequent in patients with benign primary tumors.¹

This case is reported because of the simultaneous occurrence of two distinct benign cardiac tumors. One of these we believe contributed to the production of cardiac arrhythmias.

Case report

A 43-year-old Caucasian man was seen at the Henry Ford Hospital with a twenty-year history of episodes of palpitations. Although he had had a number of medical evaluations an etiology for the palpitations was never established. Approximately three weeks prior to his referral to our hospital he was hospitalized elsewhere because of severe substernal chest pain. This radiated into the left arm and fingers and was accompanied by dyspnea and diaphoresis. The pain continued intermittently for several days. During this time serum creatine phos-

phatase (CPK), serum glutamic oxalacetic transaminase (SGOT), lactic dehydrogenase (LDH) determinations and electrocardiograms (ECG) were normal except for frequent premature ventricular contractions (PVC) and supraventricular tachycardia (Fig. 1). The initial blood pressure was 100/110 mm with subsequent blood pressure being normal. The patient and two of his brothers were considered to have borderline diabetes.

Physical examination at the Henry Ford Hospital revealed a well-developed man in no acute distress with a blood pressure of 132/88 mm Hg. The lateral border of cardiac dullness was 13 cm. from the midsternal line. P_2 was greater than A_2 , M_1 greater than M_2 and the heart rate was 88 per minute. Other findings of the physical examination were normal.

Laboratory studies. Chest fluoroscopy revealed slight cardiomegaly with no specific chamber enlargement. Fasting blood sugar was 80 mg per cent two-hour postcibal (PC) blood sugar 105 mg per cent blood urea nitrogen (BUN) 11 mg per cent urinalysis with normal limits prothrombin time 13 sec with a control of 12 sec hemoglobin 14.7 Gm per cent white blood cells (WBC) 11,500 per cubic millimeter with 2 neutrophils, 4 bands, 10 neutrophils, 2 eosinophils, 21 lymphocytes and 5 monocytes. Venereal Disease Research Laboratory (VDRL) test nonreactive triglyceride 132 mg per cent uric acid 7.1 mg per cent serum calcium 9.2 mg per cent serum phosphate 3.3 mg per cent serum complement 238.9 units total cholesterol 200 mg per cent Serum protein electrophoresis albumin 3.64 Gm per cent alpha 0.21 Gm per

From Henry Ford Hospital, Detroit, Mich.

Received for publication August 31, 1972.

Reprint requests to Daniel T. Anbe, M.D., Associate Physician, Division of Cardiac and Vascular Disease, Henry Ford Hospital, 99 W. Grand St., Detroit, Mich. 48202.

- Use of sympathetic beta-receptor blocking agents in the investigation of reflex changes in heart rate *Br J Pharmacol* 24:781 1965
- 16 Robinson B I Epstein S I Beiser G D and Braunwald D I Control of heart rate by the autonomic nervous system *Circ Res* 19:400 1966
- 17 McKenna D H Corliss R J Sailer S Zornstorf W C Crumpton C W and Rowe G G Effect of propranolol on systemic and coronary haemodynamics at rest during simulated exercise *Circ Res* 19:520 1966
- 18 Keroes J Ecker R R and Rapoport H Ventricular function curves in the exercising dog *Circ Res* 25:557 1969
- 19 Pitt H and Gregg D E Coronary haemodynamic effects of increasing ventricular rate in the unanaesthetized dog *Circ Res* 22:753 1968

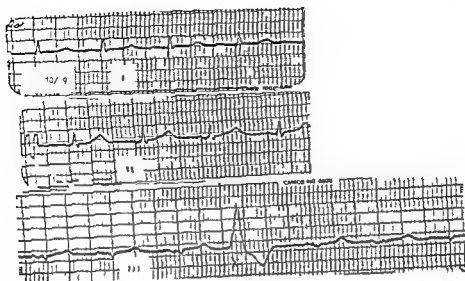


Fig 3 Leads I II III Pacemaker shifting from the sinus node to the junction best shown in Lead II

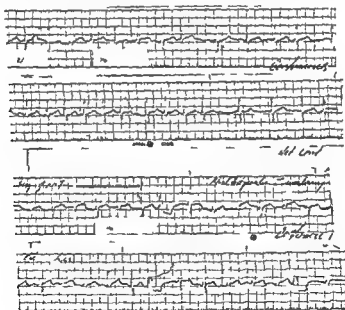


Fig 4 Monitor lead Atrial or sinus tachycardia with wandering pacemaker (inverted P wave of junctional beat)

base of the aorta the epicardium appeared gray firm and fibrous. Transsection through this area revealed a thick firm fibrous tissue containing a number of spaces 1 to 2 mm in diameter.

A discrete yellow moderately firm 2.6 by 2.1 by 3 cm nodule was present in the right atrial wall in the region of the Eustachian valve and coronary sinus. It slightly distorted these two structures and protruded into the right atrial chamber (Figs 9 and 10). Cross sections through the nodule revealed it to be

adipose tissue that was well delineated from the surrounding atrial wall. Replacement of the myocardium by adipose tissue was diffuse and marked in the right atrium, moderate in the right ventricle and minimal in the left atrium.

Microscopic pathology. The epicardium in the area of the entire A-V sulcus and the adventitia of the root of the aorta were thickened by poorly cellular fibrous tissue which contained many empty variably sized tortuous channels (Figs 11 to 13). These

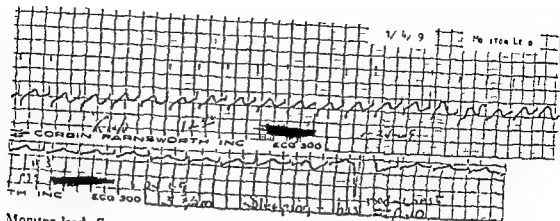


Fig 1 Monitor lead Supraventricular tachycardia suggestive of paroxysmal atrial tachycardia Occasional premature ventricular contraction

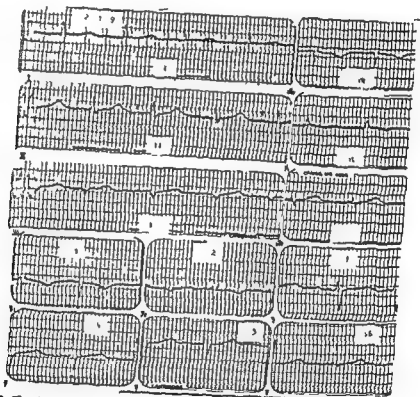


Fig 2 Twelve lead ECG Normal ECG except for a negative T wave in aVL

cent alpha₂ 0.96 Gm per cent beta globulin 1.03 Gm per cent gamma globulin 1.14 Gm per cent total protein 7.0 Gm per cent SGOT 15 units CPK 220 and 120 units LDH 485 and 505 units

Hospital course The initial ECGs revealed a basic sinus rhythm a negative T wave in aVL occasional PVCs and many atrial premature contractions with evidence of a wandering pacemaker moving from the sinus node to the A-V junctional area (Figs 2 and 3). After this four days of continuous ECG monitoring in the coronary care unit revealed evidence of a wandering pacemaker frequent premature atrial and junctional beats occasional ventricular premature beats periods of sinus tachycardia occasional short runs of supraventricular tachycardia which resembled paroxysmal atrial tachycardia (PAT) and a period of first degree A-V block (Figs 4 to 6). The patient's chest pains sub-

sided after the first hospital day. On the ninth hospital day selective right and left coronary arteriograms were performed (Figs 7 and 8). Two injections of 4 ml of Renografin 75 were made into the right coronary artery and three injections into the left coronary artery. Following the third injection of Renografin 75 into the left coronary artery there ensued hypotension bradycardia and subsequently cardiac arrest (ventricle ventricular tachycardia ventricular fibrillation). Continued attempts at resuscitation for two hours and fifty minutes were unsuccessful. Autopsy permission was obtained.

Gross pathology The only significant autopsy finding was in the heart. The heart weighed 650 grams with hypertrophy of the myocardium of both ventricles. The valves were unremarkable and the coronary arteries showed only minimal atherosclerosis. In the A-V sulcus encompassing the entire heart and



Fig 7 Normal left coronary angiogram left anterior oblique projection

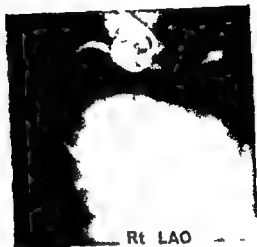


Fig 8 Normal right coronary angiogram left anterior oblique projection



Fig 9 The circumscribed tumor (A) of the interatrial septum distorts the mitral valve and coronary sinus. Adipose tissue has replaced much of the muscle of the right atrium (B), right ventricle (C), and A-V node (D).

atrial wall was an accumulation of fat cells with few interspersed cardiac muscle fibers. The myocardial fibers of the walls of all the chambers were increased in size and contained large hyperchromatic nuclei. Focal small areas of fibrosis were present in the left ventricular muscle.

Discussion

A number of diagnostic interpretations are prompted by the composite of vascular

spaces, adipose tissue, and smooth muscle diffusely distributed throughout both atrial walls and adventitia of the base of the aorta. These include (1) malformation (hamartoma) of multiple mesenchymal tissues, (2) lymphangioma associated with a marked adipose infiltration of cardiac muscle with the formation of a discrete tumor of adipose tissue of the right atrial muscle

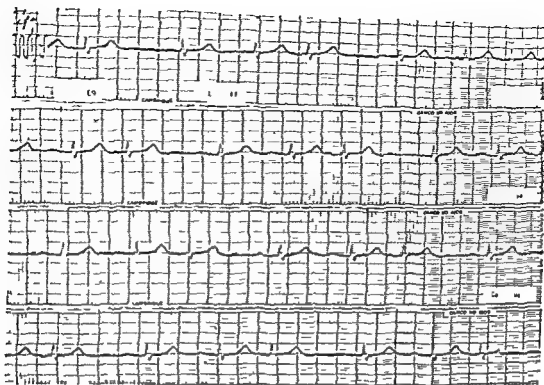


Fig 5 Long Lead II First degree A V block with PR of 0.22 sec Premature atrial contractions with atrial trigeminy Note that the atrial premature beats have a shorter PR interval of 0.16 sec

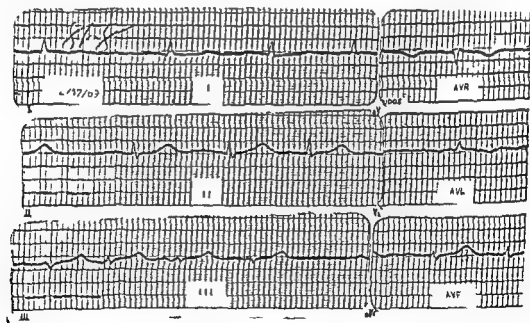


Fig 6 Junctional tachycardia in Lead III

channels were lined by uniform flat endothelial cells and had walls of thin fibrous tissue or fibrous tissue with smooth muscle that were focally thickened. Although irregularly arranged thin elastic fibers were present in some of the larger channels, no internal or external elastic lamina could be found. Similar endothelial lined channels, variously sized nerves, bundles of smooth muscle, and adipose tissue replaced most of the muscle of the right atrium and interatrial septum and, to a lesser extent, the muscle of the left atrium. Muscle replacement by adipose

tissue was diffuse and very extensive in the right atrium in the atrial septum and in the sinoatrial and A V nodes (Fig 14). It was moderate in the right ventricle and minimal in the left atrium. Distant from the A V sulcus, the epicardium of the atria and right ventricle was not fibrotic but contained increased adipose tissue. This tissue was in continuity with the fat cells in the underlying cardiac muscle and often was not accompanied by the endothelial lined channels. The polypoid nodule in the right



Fig 7 Normal left coronary angiogram left anterior oblique projection

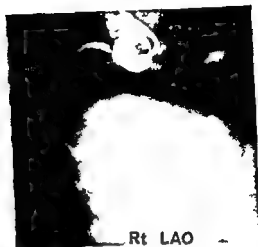


Fig 8 Normal right coronary angiogram left anterior oblique projection



Fig 9 The circumscribed tumor (A) of the interatrial septum distorts the Eustachian valve and coronary sinus. Adipose tissue has replaced much of the muscle of the right atrium (B), right ventricle (C), and A-V node (D).

atrial wall was an accumulation of fat cells with few interspersed cardiac muscle fibers. The myocardial fibers of the wall of all the chambers were increased in size and contained large hyperchromatic nuclei. Focal small areas of fibrosis were present in the left ventricular muscle.

Discussion

A number of diagnostic interpretations are prompted by the composite of vascular

spaces, adipose tissue, and smooth muscle diffusely distributed throughout both atrial walls and adventitia of the base of the aorta. These include (1) malformation (hamartoma) of multiple mesenchymal tissues, (2) lymphangioma associated with a marked adipose infiltration of cardiac muscle with the formation of a discrete tumor of adipose tissue of the right atrial muscle

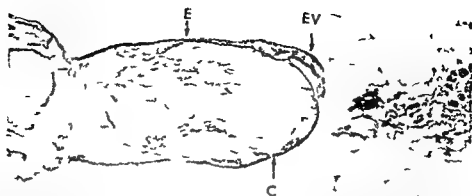


Fig. 10 (A left) Section through the tumor (1 of Fig. 9). It is differentiated adipose tissue replacing most of the cardiac muscle fibers. EV Luschka valve. E endocardium of right atrium. C endothelial surface of coronary sinus (Hematoxylin and eosin 26X). (B right) Magnification of an area of (A) (Hematoxylin and eosin 175X).

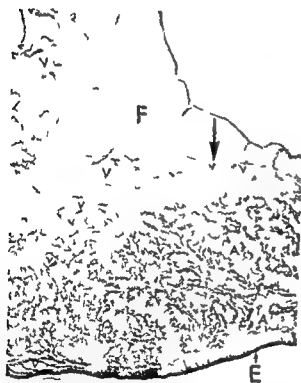


Fig. 11 Full thickness section through the left atrial wall and AV sulcus. Many empty channels (V) and adipose tissue (1) are diffusely distributed throughout the atrial muscle to the subendocardial area. Poorly cellular fibrous tissue (F) is found principally in the epicardium replacing the adipose tissue. E Left atrial endocardium (Hematoxylin and eosin 75X).

and (3) a mixed mesenchymal neoplasm (benign mesenchymoma). A lymphatic rather than a blood vascular growth is indicated by (1) the absence of a well developed muscle coat with internal and external elastic lamina in the channels that would be expected in veins and arteries of comparable size, (2) failure to demonstrate a

vascular abnormality in the coronary arteriogram (3) absence of blood in the endothelial lined channels and (4) the histologic similarity between the cystic lymphangioma (hygroma) occurring in other areas and the present growth.

Eleven cardiac tumors composed of more than one type of mesenchymal tissue—adipose, vascular and smooth muscle—have been reported.^{2,4} Depending on the authors' views regarding neoplasia and malformation the terms hamartoma, benign mesenchymoma, lymphangioma or an elaborate descriptive diagnosis have been used. Unlike the case under discussion seven of the tumors were discrete masses.⁴ Six involved the epicardium and one involved the myocardium of the right ventricular outflow tract. Four tumors were diffuse involving the left ventricular myocardium in two instances, the myocardium of both ventricles in one case and the AV node in the fourth case. In five cases the tumors did not produce symptoms and were discovered at necropsy or as a cardiac enlargement in a routine chest x-ray. Dyspnea, chest pain, systolic cardiac murmur, palpitations and pericardial effusion were present singly or in combination in four of the cases and heart block was present in one case. Successful surgical removal of the tumor was possible in five cases.

The microscopic descriptions and photomicrographs indicate that there are two tumor types among these 11 cases which differ primarily in their vascular makeup. In one group the vessels are large with a well formed muscle coat and elastic lamina



Fig 12 (A left) Area of Fig 11 (arrow) magnified (Hematoxylin and eosin 70X) (B right) Some of the smooth muscle bundles (M) not forming part of the channels are depicted (Hematoxylin and eosin 240X)



Fig 13 (A left) The adventitia (A) of the aorta (AO) is thickened by fibrous tissue which contains many lymphatics (V) in addition to the normal blood vessels and adipose tissue (Orcein azure eosin stain 13X) (B right) Area of (A) (arrow) magnified (A Adipose tissue V lymphatic channels (Orcein azure eosin stain 10X))

while in the other group they are predominantly small with little or no muscle coat. It appears that the former are best classified as arterial and/or venous malformations

(hamartomas) and the latter as lymphangiomas. The case of the present report is considered another example of diffuse lymphangioma with the added feature of

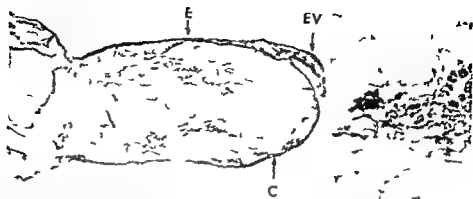


Fig. 10 (A left) Section through the tumor (1 of Fig. 9). It is differentiated adipose tissue replacing most of the cardiac muscle fibers. EV, Eustachian valve; E, endocardium of right atrium; C, endothelial surface of roof of the coronary sinus. (Hematoxylin and eosin 2.6X). (B right) Magnification of an area of (4). (Hematoxylin and eosin 175X).

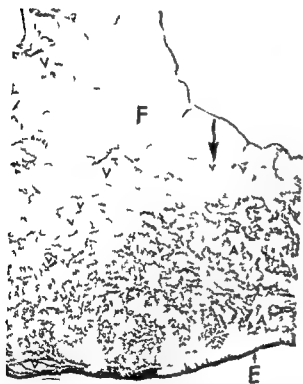


Fig. 11 Full thickness section through the left atrial wall and A V sulcus. Many empty channels (V) and adipose tissue (A) are diffusely distributed throughout the atrial muscle to the subendocardial area. Poorly cellular fibrous tissue (F) is found principally in the epicardium replacing the adipose tissue. E, Left atrial endocardium. (Hematoxylin and eosin 7.5X).

and (3) a mixed mesenchymal neoplasm (benign mesenchymoma). A lymphatic rather than a blood vascular growth is indicated by (1) the absence of a well developed muscle coat with internal and external elastic lamina in the channels that would be expected in veins and arteries of comparable size, (2) failure to demonstrate a

vascular abnormality in the coronary arteriogram (3) absence of blood in the endothelial lined channels and (4) the histologic similarity between the cystic lymphangioma (lyngroma) occurring in other areas and the present growth.

Eleven cardiac tumors composed of more than one type of mesenchymal tissue—adipose, vascular, and smooth muscle—have been reported.^{2,12} Depending on the author's views regarding neoplasia and malformation, the terms hamartoma, benign mesenchymoma, lymphangioma, or an elaborate descriptive diagnosis have been used. Unlike the case under discussion, seven of the tumors were discrete masses. Six involved the epicardium and one involved the myocardium of the right ventricular outflow tract. Four tumors were diffuse involving the left ventricular myocardium in two instances, the myocardium of both ventricles in one case, and the A V node in the fourth case. In five cases the tumors did not produce symptoms and were discovered at necropsy or as a cardiac enlargement in a routine chest x-ray. Dyspnea, chest pain, systolic cardiac murmur, palpitations, and pericardial effusion were present singly or in combination in four of the cases and heart block was present in one case. Successful surgical removal of the tumor was possible in five cases.

The microscopic descriptions and photomicrographs indicate that there are two tumor types among these 11 cases which differ primarily in their vascular makeup. In one group the vessels are large with a well formed muscle coat and elastic lamina

process associated with a lipomatous replacement of atrial cardiac muscle which in one area formed a discrete tumor a lipoma

REFERENCES

- 1 Friendberg C K. Disease of the heart Phila delphia 1966 W B Saunders Company Chap 46 p 1712
- 2 Childress R H King E D Aldrich D D Buehl J A King H and Genovese P H Successful resection of a benign right ventricular mesenchymoma Am J Cardiol 20:255 1967
- 3 Franciosi R A Gay R M and Ah Tye P Vascular hamartoma of the heart in a child Am HEART J 79:676 1970
- 4 Grant R T and Camp P D A case of complete heart block due to an arterial angioma Heart 16:137 1933
- 5 Havari V Siska K and Klein F Ueber eine mit Erfolg operierte Herzgeschwulst von interessantem feingeweblichem Aufbau Cardiologia 29:132 1956
- 6 Hochberg L A and Robinson A I Primary tumor of the pericardium involving the myocardium—surgical removal Circulation 1:805 1950
- 7 Lyburner H M Tumors of the heart Histopathological and clinical study Can Med Assoc J 30:368 1934
- 8 May I A Hardy K L Char F and Mc Adams J Vascular hamartoma of the right atrium with successful resection Ann Thorac Surg 1:64 1965
- 9 Neugebauer W Hamartom des Herzens M Allg Pathol 70:2 1938
- 10 Nicks R Hamartoma of the right ventricle J Thorac Cardiovasc Surg 47:162 1964
- 11 Pommer G Cited by Childress R H et al Successful resection of a benign right ventricular mesenchymoma Am J Cardiol 20:255 1967
- 12 Uehlinger E Die Lymphgefäßgeschwulste des Herzens Beitr Pathol Anat 78:434 1927
- 13 Page D L Lipomatous hypertrophy of the cardiac interatrial septum Hum Pathol 1:151 1970
- 14 Nuzum F Fatty infiltration (lipomatosis) of the auriculoventricular bundle of His with sudden unexpected death Arch Intern Med 134:640 1944
- 15 Spain D M and Cathcart R T Heart block caused by fat infiltration of the interventricular septum (cor adiposum) Am HEART J 32:659 1946
- 16 Ross C F and Belton E M A case of isolated cardiac lipodosis Br Heart J 30:726 1968
- 17 Balsaver A M Morales A R and Whitehouse F W Fat infiltration of the myocardium as a cause of cardiac conduction defect Am J Cardiol 19:261 1967
- 18 Kaminsky N I Killip T III Alonso D R and Hagstrom J W C Heart block and mesothelioma of the atrioventricular node Am J Cardiol 20:248 1967
- 19 Fine G and Morales A R Mesothelioma of the atrioventricular node Arch Pathol 92:402 1971

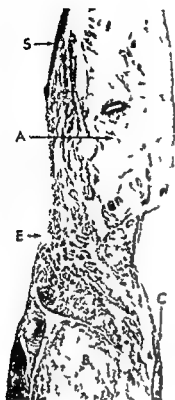


Fig 14 Adipose tissue has replaced most of the muscle of the sinoatrial node (A) and right atrial wall (B). The epicardium at the A-V sulcus is thickened by fibrous tissue (C). S Superior vena cava. E endocardium of right atrium (Hematoxylin and eosin 65X).

both a localized and diffuse lipomatous infiltration of the myocardium, the former representing a lipoma. A lymphatic vascular origin rather than a blood vascular origin is supported by the failure to visualize the tumor in the angiocardigraphic studies of one of the reported cases and by the coronary arteriogram in the present case.

The controversial question of neoplasm vs malformation for this type of growth and the relationship of the localized and diffuse lipomatous change in the cardiac muscle to the lymphangiomatous changes are not resolved by this study. The diffuse and localized lipomatous changes are not uncommon in the right atrial and atrial septal muscle and could represent a fortuitous association with the lymphangioma or an integral part of a malformation or a benign neoplasm.¹¹ The degree of lipomatous myocardial change is unusual in relation to the patient's age, normal body

weight, and the absence of a similar alteration in other organs. Its presence in the right ventricular and atrial muscle unassociated with the lymphatic channels suggests that it is independent of the lymphangioma.

Whether one considers this type of growth a hamartoma, benign mesenchymoma or a lymphangioma is less important than the recognition of its nonmetastasizing character, variability of its local behavior, diffuse epicardial and myocardial involvement, and discrete solitary tumorlike growth. Although the degree of epicardial fibrosis was unusual, it should be differentiated from a myocardial fibroma which has been occasionally designated as a hamartoma.

A casual relation between the cardiac muscle replacement and interruption of conduction system tracts and the clinical manifestations referable to the heart is strengthened by the failure to demonstrate significant coronary artery or myocardial disease by coronary arteriogram and tissue examination. Marked cardiac tissue replacement in the region of the sinoatrial node, A-V node and bundle of His has been associated with sinus bradycardia, atrial tachycardia, sinus arrests, junctional beats, atrial fibrillation, premature atrial beats and conduction abnormalities. The infiltrating tissue has variously been mesothelioma, angioma, lipoprotein and fat as in cases of lipomatous infiltration of the interatrial septum.^{4, 12, 18}

In the case presented here there was replacement and interruption of cardiac muscle in vital areas by a variety of tissues—adipose, vascular, smooth muscle and fibrous. We believe that the conduction system disturbance can be attributed to these changes.

Summary

Multiple supraventricular arrhythmias in a 43 year old man are attributed to the interruption and replacement of the specialized muscle of the conduction system. The morphologic appearance of the vascular channels and their failure to fill in the coronary arteriogram indicate that the channels are of a lymphatic origin and that the growth is one of a diffuse lymphangiomatous

process associated with a lipomatous replacement of atrial cardiac muscle which in one area formed a discrete tumor a lipoma

REFERENCES

- 1 Friendberg C K Disease of the heart Philadelphia 1966 W B Saunders Company Chap 46 p 1712
- 2 Childress R H King R D Aldrich D D Buehl J A King H and Genovese P D Successful resection of a benign right ventricular mesenchymoma Am J Cardiol 20:255 1967
- 3 Franciosi R A Gay R M and Ah Tye P Vascular hamartoma of the heart in a child Am HEART J 9 676 1970
- 4 Grant R T and Camp P D A case of complete heart block due to an arterial angioma Heart 16 137 1933
- 5 Havari V Siska K and Klein F Uber eine mit Erfolg operierte Herzgeschwulst von interessanter feingeweblichem Aufbau Cardiologia 29:132 1956
- 6 Hochberg L A and Robinson A I Primary tumor of the pericardium involving the myocardium—surgical removal Circulation 1 805 1950
- 7 Lymburner R M Tumors of the heart Histopathological and clinical study Can Med Assoc J 30:368 1934
- 8 Lay J A Hardy K L Char F and Mc Adams J Vascular hamartoma of the right atrium with successful resection Ann Thorac Surg 1 64 1965
- 9 Neugebauer W Hamartom des Herzens Z Allg Pathol 70:2 1938
- 10 Nicks R Hamartoma of the right ventricle J Thorac Cardiovasc Surg 47:762 1964
- 11 Pommer G Cited by Childress R H et al Successful resection of a benign right ventricular mesenchymoma Am J Cardiol 20:255 1967
- 12 Uehlinger E Die Lymphgefäßgeschwulste des Herzens Beitr Pathol Anat 78:434 1927
- 13 Page D L Lipomatous hypertrophy of the cardiac interatrial septum Hum Pathol 1:151 1970
- 14 Duzum F Fatty infiltration (lipomatosis) of the auriculoventricular bundle of His with sudden unexpected death Arch Intern Med 13 640 1914
- 15 Spain D M and Cathcart R T Heart block caused by fat infiltration of the interventricular septum (cor adiposum) Am HEART J 32 659 1946
- 16 Ross C F and Belton E M A case of isolated cardiac lipoidosis Br Heart J 30 726 1968
- 17 Balsaver A M Morales A R and Whitehouse F W Fat infiltration of the myocardium as a cause of cardiac conduction defect Am J Cardiol 19:261 1967
- 18 Kaminsky N I Killip T III Alonso D R and Hagstrom J W C Heart block and mesothelioma of the atrioventricular node Am J Cardiol 20 248 1967
- 19 Fine C and Morales A R Mesothelioma of the atrioventricular node Arch Pathol 92:402 1971

Propranolol induced alopecia

*Carroll M Martin MD Major MC, USA**

Eduard G Southwick, MD Major, MC USA

Howard I Maibach MD

San Francisco Calif

Propranolol hydrochloride has gained widespread clinical usage for a variety of cardiovascular disorders. It is generally a well tolerated drug but a variety of undesired effects, unrelated to its pharmacologic effects, have been observed. These include mental confusion, nausea, vomiting, diarrhea, palpitations, rashes, sleeplessness, fatigue, and light-headedness.¹ Three instances of reversible alopecia coincident with propranolol hydrochloride administration were reported during the clinical investigation of the drug.² However, direct relationship of the alopecia to the administration of the drug was not established. The following report describes an instance of reversible alopecia secondary to administration of propranolol hydrochloride in which the causal relationship is strongly suggested by drug rechallenge.

Case report

A 36-year-old man developed supraventricular tachycardia in 1959 and was diagnosed as having the Wolff-Parkinson-White syndrome in 1962. At that time he was treated with digoxin, quinidine, and procainamide. Over the ensuing years he had recurrent episodes of supraventricular tachyarrhythmias and was treated with a variety of drugs in addition

to those mentioned above including reserpine, guanethidine, hydroxazine, and diphenhydantoin. He was unresponsive to these medications as well as to induction of hypothyroidism with methimazole, stellate ganglion block, and paired atrial stimulation. In 1965 he had a two-month trial of therapy with propranolol hydrochloride in a dose ranging from 10 mg to 30 mg four times daily. Again in 1967 he had a three-month trial with propranolol in similar doses. On each occasion there was little response of his arrhythmia, but on neither occasion was alopecia noted.

In September 1970 he was again started on propranolol hydrochloride 10 mg twice daily. Additional medications at that time included gitalin and quinidine. Over the next three months the dose of propranolol hydrochloride was increased to 40 mg four times daily. In January 1971 he noted for the first time a patchy hair loss involving his scalp, chest, and arms. This alopecia progressed slowly until May 1971 when the process seemed to suddenly accelerate. In June 1971 he consulted his physician and the propranolol hydrochloride was discontinued. At that time there were patchy areas of hair loss over the scalp and arms (Fig. 1) and centrally diffuse hair loss from the chest (Fig. 2). The underlying skin appeared normal.

Telogen counts from the periphery of the bald areas of the scalp averaged 68 per cent (normal scalp averages 15 per cent telogen hairs³ and telogen counts over 25 per cent are diagnostic of telogen effluvium⁴). A biopsy of the scalp in an area of alopecia was normal except for the increased number of telogen hairs.

Microscopic examination of scalp and body hairs

From the Cardiology Service and Dermatology Service, Letterman General Hospital, San Francisco, Calif., and the Department of Dermatology, University of California Medical Center, San Francisco, Calif.

Received for publication Sept. 18, 1972.

Reprint requests to: Technical Publications Editor, Letterman General Hospital, San Francisco, Calif. 94129.

*Present affiliation: Division of Cardiology, Department of Medicine, Madigan General Hospital, Tacoma, Wash.

†Harvey A. Barnett, MD, Assistant Medical Director, Ayerst Laboratories, New York, N.Y., in addition to Doctor Martin dated October 11, 1971.



Fig. 1 The patient's arm showing diffuse patchy alopecia.

removed by gently pulling hairs from the periphery of the bald patches revealed normal appearing telogen and anagen hairs. However, with polarized light microscopy, the telogen hair bulbs were noted to be dysplastic and there was a dysplastic defect in each of these hair shafts about one third distal from the bulb. Anagen hairs appeared normal under polarized light microscopy.

The alopecia progressed over the next two months until he had extensive hair loss from his chest, forearms, and eyebrows and approximately 50 per cent loss from his scalp. By the end of the fourth month after discontinuing propranolol hydrochloride he had almost complete resolution of the alopecia. At this time after informed consent had been obtained he was re-challenged with propranolol hydrochloride 80 mg four times daily. Within four weeks he had definite recurrence of the alopecia, again beginning on the scalp, chest, and forearms. The shed hairs were telogen. Propranolol hydrochloride was again discontinued and over the next four months he had gradual and complete return of his hair growth.

Comment

Drug induced alopecia is characteristically the anagen effluvium type. The drug acts on the hair follicle to inhibit mitosis. Examples of drugs that cause this type of alopecia are the antimetabolic agents, folic



Fig. 2 The patient's chest showing a central more diffuse hair loss.

acid antagonists, purine antagonists, alkylating agents, thallium, and colchicine.

Telogen effluvium following drug therapy is rarely reported except for heparin induced alopecia. This is an unusual case of telogen effluvium with dysplasia of the hair bulb and hair shaft caused by propranolol.

The criteria of Kligman³ for telogen effluvium—i.e. (1) huge telogen counts—25 per cent to 60 per cent (2) latency period of about 3 months between stress and shedding (3) loss of club hairs from normal telogen follicles and (4) regrowth with drug cessation have been fulfilled. Also when re-challenged with the drug there was recurrence of hair loss. The re-challenge showed that the drug induced telogen hair loss occurred more rapidly (one month) than that seen with stress induced (fever etc.) telogen effluvium.

Summary

A patient developed diffuse patchy alopecia following propranolol therapy. This was reproduced by re-challenge with the drug. The alopecia demonstrated the criteria for telogen effluvium.

REFERENCES

1. Stephen S A. Unwanted effects of propranolol. *Am J Cardiol* 33:163, 1966.
2. Van Scott F J, Reinertson R I, and Steinmuller R. The growing hair roots of the human scalp and morphologic changes therein following amethopterin therapy. *J Invest Dermatol* 29:197, 1957.
3. Kligman A M. Pathologic dynamics of human hair loss. *Arch Dermatol* 83:175, 1961.

Propranolol induced alopecia

*Carroll M Martin MD, Major, MC USA**

Edward G Southwick MD Major MC, USA

Howard I Warbach, MD

San Francisco, Calif

Propranolol hydrochloride has gained widespread clinical usage for a variety of cardiovascular disorders. It is generally a well tolerated drug but a variety of undesired effects unrelated to its pharmacologic effects have been observed. These include mental confusion, nausea, vomiting, diarrhea, palpitations, rashes, sleeplessness, fatigue, and light headedness.¹ Three instances of reversible alopecia coincident with propranolol hydrochloride administration were reported during the clinical investigation of the drug.* However, direct relationship of the alopecia to the administration of the drug was not established. The following report describes an instance of reversible alopecia secondary to administration of propranolol hydrochloride in which the causal relationship is strongly suggested by drug rechallenge.

Case report

A 36 year old man developed supraventricular tachycardia in 1959 and was diagnosed as having the Wolff Parkinson White syndrome. At that time he was treated with digoxin, quinidine, and procainamide. Over the ensuing years he had recurrent episodes of supraventricular tachyarrhythmias and was treated with a variety of drugs in addition

to those mentioned above including reserpine, guanethidine, hydroxyzine, and diphenylhydantoin. He was unresponsive to these medications, as well as to induction of hypothyroidism with methimazole, stellate ganglion block, and paired atrial stimulation. In 1965 he had a two-month trial of therapy with propranolol hydrochloride in a dose ranging from 10 mg to 30 mg four times daily. Again in 1967 he had a three month trial with propranolol in similar doses. On each occasion there was little response of his arrhythmia, but on neither occasion was alopecia noted.

In September 1970 he was again started on propranolol hydrochloride 10 mg twice daily. Additional medications at that time included gitalin and quinidine. Over the next three months the dose of propranolol hydrochloride was increased to 40 mg four times daily. In January 1971 he noted for the first time a patchy hair loss involving his scalp, chest, and arms. This alopecia progressed slowly until May 1971 when the process seemed to suddenly accelerate. In June 1971 he consulted his physician and the propranolol hydrochloride was discontinued. At that time there were patchy areas of hair loss over the scalp and arms (Fig 1) and centrally diffuse hair loss from the chest (Fig 2). The underlying skin appeared normal.

Telogen counts from the periphery of the bald areas of the scalp averaged 68 per cent (normal scalp averages 15 per cent telogen hairs² and telogen counts over 25 per cent are diagnostic of telogen effluvium³). A biopsy of the scalp in an area of alopecia was normal except for the increased number of telogen hairs.

Microscopic examination of scalp and body hairs

From the Cardiology Service and Dermatology Service, Letterman General Hospital, San Francisco, California, and the Department of Dermatology, University of California Medical Center, San Francisco, California.

Received for publication Sept. 18, 1972.

Reprint requests to: Technical Publications Editor, Letterman General Hospital, San Francisco, California 94129.

Present affiliation: Division of Cardiology, Department of Medicine, Madigan General Hospital, Tacoma, Washington.

**Harvey A. Barnett, MD, Assistant Medical Director, Ayer Laboratories, New Britain, Connecticut, to Doctor Martin dated October 11, 1971.*

August 1973 Vol 86

p. 1 m. 86
v. m. 82

Table I Patient data on July 9 1971

| Time (A.M.) | Location | Systolic pressure | Diastolic pressure | Mean | Rhythm | Heart rate |
|-------------|-------------|-------------------|--------------------|------|--------|------------|
| 9 48 | AO | 115 | 73 | 90 | AF | 110 |
| | PA | 63 | 26 | 41 | | |
| 10 10 | Hypotension | | | | | |
| 10 29 | AO | 33 | 70 | 97 | AF | 110 |
| | PA | 61 | 24 | 38 | | |
| | RA | C = 17 | V = 22 | 14 | | |
| 10 44 | LV | 107 | 16 20 | 29 | | |
| | RPCW | | | 28 | | |
| | RP4 | 35 | 24 | 19 | | |
| 10 47 | RA | | | | | |
| | LV | 102 | 15 20 | 74 | | |
| | AO | 100 | 71 | 29 | | |
| | PA | 35 | 25 | | | |
| | RV | 35 | 15 20 | 30 | | |
| | RA | | | | | |

Abbreviations: AO = aorta PA = pulmonary artery RA = right atrium RV = right ventricle LV = left ventricle RPCW = right pericardial catheter RP4 = right pulmonary artery C = C wave V = V wave AF = atrial fibrillation

the patient lying in bed propped up at a 30 degree angle. The chest expanded fully and symmetrically and an occasional wheeze was heard. Coarse rales were noted at the left base and minimal rales were noted at the right base. The peripheral pulses were equal in intensity and graded as 2+ for the carotids, radials and femorals. Dorsal pedal pulses were not palpable bilaterally. Examination of the heart revealed an irregular rhythm with a heart rate of 120 per minute. Paradoxical pulsations were noted over the apex. There was a loud S3. A Grade III/VI systolic murmur and a Grade II/VI mid-diastolic rumble were heard at the apex. The holosystolic murmurs radiated into the axilla. The liver and spleen were not palpable. A large reducible inguinal hernia was seen on the right. Rectal examination was within normal limits. Neurological examination was unremarkable. Examination of the extremities revealed 4+ pitting edema over the feet and tibiae.

Laboratory data and hospital course. Chest x ray was read as showing marked cardiomegaly, predominantly left atrial enlargement with clear lung fields showing no active infiltration. ECG revealed abnormal findings due to atrial fibrillation, non-specific ST-T wave changes and an occasional aberrantly conducted beat. Vectorcardiography was reported to be abnormal due to left ventricular hypertrophy with probable right ventricular hypertrophy and abnormal T waves probably secondary to left ventricular hypertrophy. Urinalysis on admission was within normal limits. Hemoglobin was 14.6 Gm per 100 cc, hematocrit was 47.5 per cent, white blood cell count was 9,400 per cubic millimeter with 80 per cent neutrophils, 19 per cent lymphocytes, 1 per cent monocytes, adequate platelets with a few atypical cells and macrocytes and a few bizarre target cells. SMA-6 was within normal limits except for a glucose of 163 mg per cent, prothrombin time was 14.4 seconds. Venereal Disease Research Laboratory test was non-reactive. On July 9, the patient was sent for

cardiac catheterization. During this procedure as the trans-septal puncture was attempted the patient became hypotensive and dyspneic. The hypotensive episode lasted from 10:00 A.M. to 10:20 A.M. with mean aortic pressure ranging from 70 to 75 mm mercury. Following this pressures from the aortic root, pulmonary artery and right atrium were again recorded. Because of progressive dyspnea of the patient the study was interrupted. However the following information (Table I) was recorded.

Following return to the ward the patient remained hypotensive with a systolic pressure of 15 to 80 mm Hg. It was noted that there was no urine output and that the urinary bladder was not distended. A Foley catheter was passed but no urine was drained. At 10:45 P.M. on July 9 1971 the patient was transferred to the Intensive Care Unit and an intra arterial line was placed in the left arm. A central venous pressure line (CVP) was placed in the right arm. At the time of admission to the Intensive Care Unit the blood pressure was 100/80 pulse rate 120 per minute. Wheezes were noted in the anterior lung fields and rales were observed in the posterior lung field. The liver was noted to be 4 fingerbreadths below the costal margin in the mid-clavicular line. The spleen was not palpable. Levothroid was administered by drip until the blood pressure was stabilized. The laboratory tests were reported as sodium 139 mEq per liter, potassium 5.6 mEq per liter, chlorides 99 mEq per liter, carbon dioxide measured 17 blood urea nitrogen 39 mg per cent, glucose 130 mg per cent, creatinine 2.9 mg per cent, calcium 5.2 mEq per liter. The nephrologist in consultation thought that the patient had acute tubular necrosis. He had no response to large doses of Lasix and was treated for renal failure with furosemide.

On the evening of July 10 1971 it was noted that the patient had large ecchymoses at several venous puncture sites and at the site of the right brachial

Clinical pathologic conference

Vahe A. Karachorlu MD*
Rolf M. Gunnar, MD**
Cecil A. Krakower MD***
Chicago Ill

Case report

Clinical abstract This was the first admission to the University of Illinois Hospital for this 49 year old Caucasian man. He was referred from an outside hospital with a diagnosis of arteriosclerotic heart disease and left ventricular aneurysm.

The patient first complained of dyspnea and weakness of sudden onset in 1956. He was hospitalized at that time and placed on digitalis and "water pills." He was then asymptomatic until 1963 when he was again hospitalized for retention of fluid. No further information could be obtained with respect to these two hospitalizations or his condition subsequent to the second one. However it is known that he was hospitalized for a third time on June 7, 1966 because of considerable increase in dyspnea and orthopnea. There had been no peripheral edema or rales. The patient was diagnosed as having arteriosclerotic heart disease with ventricular fibrillation and congestive failure. He was given diuretics and placed on digitalis. He was discharged on June 17, 1966 greatly improved and in cardiac compensation.

The patient's next admission, the fourth hospitalization since 1966, was on June 4, 1971 due to nausea, vomiting and weight loss for one week. He had been maintained on Lanoxin and Hygroton.

Physical examination at the outside hospital revealed pulse 100, cardiac respirations 22, blood pressure 130/82. There was an irregular rhythm. A 2 was louder than P 2. There were no particular murmurs or extrasystoles. The liver was one finger breadth below the right costal margin. An electrocardiogram (ECG) was reported to have shown ventricular fibrillation and incomplete left bundle branch block.

X-ray study of the chest was reported by the radiologist as follows: Preliminary fluoroscopy of the chest shows a markedly enlarged heart. Along

the left heart border there are two densities. One the main density pulsates as the heart beats normally. The other density which is anterior to it also has a rounded contour, is separate from the main density and is not as dense. It probably represents a large aneurysm along the anterior ventricular wall covering a distance of at least 6 by 7 cm. In addition to this, on barium swallow, the esophagus is deviated markedly to the right and is elevated at the level of the arch of the aorta in a manner typical of an enlargement of the left auricle. So this patient must have some type of mitral valvular defect besides the aneurysm to cause the marked enlargement of the left auricle.

For this reason the patient was transferred to the University of Illinois Hospital for further evaluation and treatment on June 25, 1971. The following pertinent information was elicited. He never had chest pain, palpitation or hemoptysis. He gave no history which could be interpreted as that of rheumatic fever but he did give a history of gouty arthritis. He had smoked one pack of cigarettes a day for the past 45 years.

He had had an appendectomy in 1943, herniorrhaphy in 1948 and cholecystectomy in 1955.

Physical examination Examination revealed a healthy appearing elderly Caucasian man in no acute distress. Blood pressure was 130/80 mm mercury. Pulse rate was irregularly irregular ranging from 90 to 130 per minute. Temperature was 98.6 F. The skin was flaccid and dry with seborrheic keratosis about the face. The head was normocephalic. The pupils were equal, round and reactive to light and accommodation. Funduscopic examination revealed mild arteriovenous nicking, copper wire changes and arteriolar narrowing. The tympanic membranes were clear. There was a mild hearing deficit. Examination of the neck revealed distended veins for a distance of 4 cm above the clavicle with

From the Department of Pathology, University of Illinois Hospital, Chicago, Ill.
Received for publication Oct. 26, 1972.

Reprint requests to Dr. Cecil A. Krakower, Head, Department of Pathology, University of Illinois at the Medical Center, 1853 W. Polk St., Chicago, Ill. 60612.

*Department of Pathology, University of Illinois Hospital, Chicago, Ill.

**Department of Medicine, Stritch School of Medicine, Chicago, Ill.

***Department of Pathology, University of Illinois Hospital, Chicago, Ill.

August 1973, Vol 86

1 me 86
1 me 2

The physical examination now done at the University hospital reveals a normal blood pressure and an irregular pulse, retinal changes which would be consistent with either arteriosclerosis or pre-existing hypertension. The neck veins were distended consistent with the diagnosis of cardiac failure but the pulsations are not described. It would be interesting to know if there were large systolic pulsations or whether the neck veins were distended and without pulsations. The former would suggest the presence of tricuspid insufficiency; the latter might suggest obstruction to filling of the right heart either by a lesion in the mediastinum, pericardial effusion or loss of compliance of the right ventricle. The absence of pedal pulses is further evidence of the patient's generalized arteriosclerosis. We are now told that there is paradoxical pulsation over the apex of the heart and this I assume means that there are two impulses pulsating asynchronously as one would see with a ventricular aneurysm. Up till this point we are eventually told that the patient has a ventricular aneurysm and the findings so far are consistent with this diagnosis. I should point out that other diagnoses are possible such as pericardial cyst or cardiac tumor to explain the x-ray shadow which has been described. Pericardial cysts usually appear at the right lower cardiac border although they can appear anteriorly and they can be thought to pulsate paradoxically. Cardiac tumors are less frequent in this age group. Left atrial myxoma has to be considered. It can produce mitral insufficiency. However, in the absence of embolic phenomena and fainting spells due to sudden episodes of obstruction I would be inclined to exclude that diagnosis. I mention myxoma not only because the obscure becomes the common in this CPC exercise but also because the patient has mitral insufficiency and a murmur described in this hospital which was not described in the other hospital immediately before transfer. Mitral insufficiency of course can cause a left lower sternal border lift and with this in addition to the apical pulsation may give the appearance of paradoxical motion. The left lower sternal lift is due to the regurgitant flow filling the left atrium, ballooning the left atrium thus pushing the heart forward against the an-

terior chest wall. In this hospital examination of the heart revealed a loud third sound Grade 3/4 systolic murmur and a Grade 2/6 mid-diastolic rumble at the apex. I assume since the murmur is at the apex that the systolic murmur is holosystolic or plateau in character and that the third sound is coming from the left ventricle. This all goes with the diagnosis of mitral insufficiency but also goes with the diagnosis of ventricular aneurysm with or without mitral insufficiency. The systolic murmur described in ventricular aneurysm is usually due to a degree of mitral insufficiency but this does not have to be severe and the mid-diastolic rumble merely represents the refilling of the left ventricular cavity from the aneurysm as well as the left atrium. The loud third sound is evidence of volume overload at the left ventricle. We now have physical findings to explain the failure either on the basis of a ventricular aneurysm or on the basis of mitral insufficiency. The mid-diastolic rumble could represent the filling rumble of severe mitral insufficiency or could represent the rumble appearing in ventricular aneurysm. Also a loud third sound consistent with either severe mitral insufficiency or ventricular aneurysm.

The liver and spleen are not palpable. This makes me a little uncomfortable as to what has happened to the liver. The patient has rather severe pitting edema of his lower extremities. I would expect that unless the patient has cirrhosis that his liver should be enlarged.

We are now treated to further information from the ECG laboratory. The vector cardiogram is reported showing left ventricular hypertrophy but there is no mention of a myocardial infarct. This makes me extremely uncomfortable with the diagnosis of ventricular aneurysm. In the absence of bundle branch block, ventricular aneurysms are almost always associated with evidence of previous myocardial infarction. This is particularly true if the aneurysm lies on the anterior wall which is the description which has been given to me so far. The rest of the laboratory examination does not particularly help me except for the fact that there is evidence of diabetes in one elevated glucose determination. This of course would be even more sup-

Table II Patient data on July 5 and on July 10 1971

| Date | Hemoglobin (Gm per cent) | Hematocrit (volumes per cent) | White blood count (per mm ³) | Platelets (per mm ³) | Prothrombin (sec) |
|---------|-----------------------------|-------------------------------------|--|-------------------------------------|----------------------|
| 7 5 71 | 14 6 | 44 5 | 9 400 | Adequate | 11 4 |
| 7 10 71 | 11 6 | 33 7 | 24 300 | 74 000 | 25 8 |

cutdown Repeat and previous hematologic examinations revealed the following (Table II) findings

Fibrinogen was 201 mg per cent Split fibrin products were positive at 1 to 16 Partial thromboplastin time was 72.8 seconds The hematologist in consultation felt that the patient possibly had disseminated intravascular coagulation and recommended treatment with heparin should the patient continue to deteriorate On July 11 1971 it was noted that the patient had a low grade fever A smear of the sputum revealed gram positive diplococci The patient was then started on procaine penicillin 600 000 units intramuscularly every 12 hours It was necessary intermittently to stabilize the patient's blood pressure with Levophed Blood cultures drawn at the time of the elevated temperature were reported positive for gram positive cocci which were later identified as *Staphylococcus aureus* sensitive to penicillin G The patient accordingly was placed on nafcillin On July 12 1971 the patient remained oliguric with a temperature of 102° F and blood urea nitrogen of 84 mg per cent On July 13 1971 heparin therapy was begun with 10 000 units stat and 7 500 units every 4 hours The patient continued to deteriorate with periods of apnea and persistent coma Chest x ray revealed an infiltrate in the left lung He was intubated and placed on an MA 1 respirator He had an episode of nodal rhythm which was attributed to digitalis toxicity Digitalis was then withheld On July 14 1971 the patient could no longer breathe spontaneously without the assistance of a respirator Blood pressure was 34/20 pulse 70 at 8 15 M All vital signs ceased at 8 45 P M

Discussion

DR GUNNAR The patient is a 79 year old man who was referred from another hospital with the diagnosis of arteriosclerotic heart disease and left ventricular aneurysm He first had evidence of cardiac failure in 1936 and this responded quite well to usual modes of treatment This episode was of sudden onset but unassociated with chest pain The patient did well until 1963 when he had another episode of failure, again treated successfully In 1956 he had evidence of left heart failure and at this time atrial fibrillation was first noted His final admission was June, 1971, with evidence of nausea, vomiting and weight loss This

was attributed to digitalis and diuretic therapy

The description of the physical examination at the hospital before transfer to the University hospital shows a normal blood pressure evidence of atrial fibrillation and some evidence of cardiac failure The ECG at that time was described as showing in complete left bundle branch block which could mean several things One could assume that the ECG loop would be such that it would produce a loss of q waves over the left precordial leads and slurring of the upstroke of the left precordial leads This could be produced by left bundle block which has no initial rightward force The initial force quickly moves posteriorly and the loop moves in a clockwise direction in a horizontal plane A similar pattern could be produced by severe left ventricular hypertrophy where the loop assumes a figure of eight configuration with the initial rightward septal forces being obliterated by the strong forces from the posterior basal portion of the left ventricle The third possibility could be anterolateral infarction where the initial anterior forces may go clockwise initially and the entire loop rotates clockwise Which of these pertain may become apparent when we see particularly the vectorcardiogram

X ray studies done at the other hospital showed an enlarged heart and a very interesting description of a double density over the anterior wall of the left ventricle as well as evidence of left atrial enlargement The conclusion from the fluoroscopy was that the patient had mitral valve disease as well as a ventricular aneurysm We are given the additional information that the patient had a past history of gouty arthritis and a long history of smoking Both of these are risk factors for the appearance of arteriosclerotic heart disease

the fibrinogen level is at the lower limits of normal the split products were positive and the partial thromboplastin time was prolonged. All of the evidence would be consistent with the diagnosis of disseminated intravascular coagulation and this was the diagnosis made by the hematologist who recommended treatment with heparin. This is indeed a difficult problem since we have a situation in which we have made the diagnosis that the patient had had a perforation of the heart at the time of catheterization with bleeding of the pericardium and now two days later because of diffuse intravascular coagulation heparin is needed to block the process. This is obviously the operation of Murphy's law which states that when anything can go wrong it will, but I am afraid that with this patient no matter what decision is made it will be wrong. I have of course the distinct advantage over the physicians treating this patient in that I know that the presentation of this patient in this amphitheater indicates that the therapy instituted was certainly not classified as a success.

Cranial negative bacteremia is much more prone to cause the diffuse intravascular coagulation since endotoxin release causes activation of Hagemann factor which in turn activates the coagulation cascade as well as activating the kinin system to cause vasodilatation and rather early cardiovascular collapse. On July 11 the patient had a low grade fever and although the sputum revealed pneumococci a blood culture was positive for *Staphylococcus aureus*. He was treated with penicillin G and later nafcillin. We are not told whether or not this was a single culture or whether the staphylococcus was coagulase positive. If this was a repeated positive culture for *Staphylococcus aureus* coagulase positive I would postulate (1) that it could have been induced by the cardiac catheter even though *Staphylococcus aureus* can be introduced by urinary catheter and (2) that there was some mediastinal abscess which had been disturbed during the time of transseptal catheterization. It appears that the patient continued to do poorly and died five days after the catheterization had been performed. I now have the task of making a final conclusion. I think I must

make two diagnoses. One for the situation prior to and the other for the condition after cardiac catheterization. I would think that the most likely diagnoses prior to catheterization are arteriosclerotic heart disease, previous myocardial infarction, mitral insufficiency perhaps due to papillary muscle dysfunction or even disruption of the chordae tendineae and less likely although possibly a ventricular aneurysm. My discomfort with the diagnosis of ventricular aneurysm is based on the lack of ECG information. My diagnosis after catheterization would be that the patient had pericardial tamponade secondary to perforation of the atrium at the time of transseptal catheterization and that he developed gram negative bacteremia and diffuse intravascular coagulation subsequent to insertion of a Foley catheter. The only way I can make a unified diagnosis would be to state that the patient did have a ventricular aneurysm and that the staphylococcal bacteremia was pre-existent to his final hospitalization that he had endocarditis involving the mitral valve thus producing mitral insufficiency and that the resurgence of his bacteremia caused diffuse intravascular coagulation and his ultimate death.

DR HOFFMAN: * The chest film taken in the outside hospital prior to the present admission reveals a transverse cardiac diameter of 19.5 cm. The heart shows generalized enlargement based on the single PA film. The lung fields show pulmonary venous hypertension with some redistribution of blood flow to the upper lung fields. There are faint Kerley lines. The views of the heart with bismum in the esophagus indicate enlargement of at least the left atrium and ventricle.

Six additional films were obtained during the present admission. Cardiac catheterization was performed on July 9, 1971. On this day the transverse cardiac diameter was 22.0 cm. This apparent increase in the silhouette can be explained by increased magnification when films are taken enteroposteriorly with the patient in the supine position. A central venous catheter was within the superior vena cava and marked enlargement of the left atrium was made apparent by considerable elevation of the

portive of the diagnosis of ventricular aneurysm since it is more common in the diabetic where silent infarction is more common. It is not uncommon to have aneurysms appear in patients who do not have documented episodes of myocardial infarction. Cardiac catheterization was performed on July 9 and the results of the catheterization are presented in the Table (Table 1). The patient had an episode of hypotension shortly after an attempt was made to cross the atrial septum with a needle. The blood pressure fell to 70 to 75 mm Hg, but after the needle was withdrawn the left heart systolic pressure returned to near normal levels. It is unfortunate that we do not have a measurement of the right atrial pressure before the episode of hypotension. The most common complication of trans septal catheterization of the left atrium is perforation of either the right or left atrium into the free pericardial cavity and occasionally there is puncture of the root of the aorta since the aorta lies in the upper portion of the atrium. As we note the right ventricular systolic pressure fell and the mean left atrial pressure went from 14 to 20. The end diastolic pressures in the left ventricle and in the right ventricle were also 20 and these findings would be consistent with pericardial effusion and tamponade. Since all four chambers of the heart are within the distended tense pericardium the end diastolic pressures in all four chambers tend to equalize. There evidently was no attempt to do pericardiocentesis and the patient was returned to the ward still somewhat hypotensive. After transfer to the Intensive Care Unit, the blood pressure was stabilized at 100/80 with a rapid pulse rate of 120. There were wheezes noted in the lungs. The liver was described as four fingerbreadths below the costal margin. The patient required Norepinephrine drip to stabilize his blood pressure but he was gradually taken off this medication. Norepinephrine may not be the agent of choice in patients with pericardial tamponade even for a short period of time to stabilize their blood pressure since this agent tends to increase the end diastolic volume. An agent which would tend to increase output at a smaller diastolic volume might be a more effective agent. The best agent for

this would be isoproterenol, but in a patient who is 76 years old with coronary artery disease there is a risk in using isoproterenol since it may lower perfusion pressure of the coronary arteries and increase myocardial ischemia. The fact is, if the patient is hypotensive because of pericardial tamponade, then the treatment does not lie in the use of any of these agents but rather in aspiration of the pericardial effusion or pericardial blood. It is somewhat difficult to aspirate much blood from these patients since the bleeding is rather vigorous and clotting occurs rather promptly within the pericardial cavity. However, removing small amounts of fluid may relieve the tamponade sufficiently to allow time to decide whether or not surgical evacuation is or is not the treatment of choice. If there is a good deal of blood in the pericardial cavity, this in my opinion, should be evacuated since this is an excellent stimulant for the formation of pericardial fibrosis and later development of constrictive pericarditis. Subsequent to the episode of hypotension the patient developed renal failure thought to be due to acute tubular necrosis. On July 10 the patient was noted to have ecchymoses at several puncture sites and at the site of the cutdown. This is one day after the catheterization. At this point there is a decrease in hemoglobin noted and although elevation of temperature is not recorded there is an elevation of white count to 24,300, decrease in the platelets to 74,000 and a decrease in the prothrombin time to 25.8 seconds. All of these findings could be due to diffuse intravascular coagulation and the question arises as to the etiology. Of course the most common reason is sepsis and the obvious source would be the cardiac catheterization. However, this is a very rare complication of cardiac catheterization so those of us who are familiar with this feature would look for other sites of sepsis. My inclination would be to look at another catheter—the Foley catheter which was inserted into the bladder after the patient developed his shock like state. This is a common source of gram negative sepsis and as we demonstrated at this institution, urinary tract instrumentation is associated with gram negative bacteremia in about 30 per cent of the patients. We feel that

J. I. M. 86
N. M. B. 12

the fibrinogen level is at the lower limits of normal, the split products were positive and the partial thromboplastin time was prolonged. All of these would be consistent with the diagnosis of disseminated intravascular coagulation and this was the diagnosis made by the hematologist who recommended treatment with heparin. This is indeed a difficult problem since we have a situation in which we have made the diagnosis that the patient had had a perforation of the heart at the time of catheterization with bleeding of the pericardium and now two days later because of diffuse intravascular coagulation heparin is needed to block the process. This is obviously the operation of Murphy's law which states that when anything can go wrong it will, but I am afraid that with this patient no matter what decision is made it will be wrong. I have of course the distinct advantage over the physicians treating this patient in that I know that the presentation of this patient in this amphitheater indicates that the therapy instituted was certainly not classified as a success.

Gram negative bacteremia is much more prone to cause the diffuse intravascular coagulation since endotoxin release causes activation of Hagemann factor which in turn activates the coagulation cascade as well as activating the kinin system to cause vasodilatation and rather early cardiovascular collapse. On July 11 the patient had a low grade fever and although the sputum revealed pneumococci a blood culture was positive for *Staphylococcus aureus*. He was treated with penicillin G and later nafcillin. We are not told whether or not this was a single culture or whether the staphylococcus was coagulase positive. If this was a repeated positive culture for *Staphylococcus aureus* coagulase positive I would postulate (1) that it could have been induced by the cardiac catheter even though *Staphylococcus aureus* can be introduced by urinary catheter and (2) that there was some mediastinal abscess which had been disturbed during the time of transseptal catheterization. It appears that the patient continued to do poorly and died five days after the catheterization had been performed. I now have the task of making a final conclusion. I think I must

make two diagnoses. One for the situation prior to and the other for the condition after cardiac catheterization. I would think that the most likely diagnoses prior to catheterization are arteriosclerotic heart disease, previous myocardial infarction, mitral insufficiency perhaps due to papillary muscle dysfunction or even disruption of the chordae tendineae and less likely although possibly a ventricular aneurysm. My discomfort with the diagnosis of ventricular aneurysm is based on the lack of ECG information. My diagnosis after catheterization would be that the patient had pericardial tamponade secondary to perforation of the atrium at the time of transseptal catheterization and that he developed gram negative bacteremia and diffuse intravascular coagulation subsequent to insertion of a Foley catheter. The only way I can make a unified diagnosis would be to state that the patient did have a ventricular aneurysm and that the staphylococcal bacteremia was pre-existent to his final hospitalization that he had endocarditis involving the mitral valve thus producing mitral insufficiency and that the resurgence of his bacteremia caused diffuse intravascular coagulation and his ultimate death.

DR. HOFFMAN: * The chest film taken in the outside hospital prior to the present admission reveals a transverse cardiac diameter of 19.5 cm. The heart shows generalized enlargement based on the single PA film. The lung fields show pulmonary venous hypertension with some redistribution of blood flow to the upper lung fields. There are faint Kerley lines. The view of the heart with barium in the esophagus indicates enlargement of at least the left atrium and ventricle.

Six additional films were obtained during the present admission. Cardiac catheterization was performed on July 9, 1971. On this day the transverse cardiac diameter was 22.0 cm. This apparent increase in the silhouette can be explained by increased magnification when films are taken anteroposteriorly with the patient in the supine position. A central venous catheter was within the superior vena cava and marked enlargement of the left atrium was made apparent by considerable elevation of the



Fig. 1 The tricuspid leaflets show multiple recent vegetations on the valve and chordae tendineae (arrows)



Fig. 2 The mitral valve has minimal fibrotic and atherosclerotic changes. Note the old ruptured chorda tendinea with thickened end (arrow)

left main stem bronchus. On July 12, 1971, the transverse cardiac diameter was 23.75 cm, which represents an increase in silhouette size when one allows for one centimeter difference between systole and diastole. There was blunting of the left costophrenic angle, apparently by fluid.

The cardiac size in the final two films, the evening of July 12, 1971, and on the day of death on July 18, 1971, could not be evaluated, since the left heart border was obscured by considerable fluid within

the left hemithorax. There was also fluid within the right hemithorax. Without the additional aid of isotope scans, ultrasonograms, or angiocardiograms, it is not possible to distinguish between cardiac enlargement or pericardial effusion in accounting for the widened cardiac silhouette.

DR KARACHORLU. At autopsy there was 2+ edema of both legs. There was a decubitus ulcer of the right buttock measuring 10 cm in diameter.

The right pleural cavity contained 440

J. J. me 86
V. mb 2



Fig 3 The posterior wall of the left ventricular epicardium in the course of the posterior descending branch is discolored and roughened. This area is outlined by arrows.

c.c. and the left 370 c.c. of clear yellowish fluid. The peritoneal cavity contained 300 c.c. of similar fluid. The pericardial sac was extremely distended with 500 c.c. of fresh blood. The heart was markedly enlarged *in situ*. It weighed 600 grams. There was no exudate over the epicardium. There was, however, a dark reddish brownish discolored roughened area over the posterior wall of the left ventricle near the apex measuring 2 by 2 cm. No perforation could be identified in this area. The right atrium was moderately enlarged measuring 8 cm. from the superior to inferior vena cava and 14 cm. in circumference. Its endocardium was normal. No hemorrhage or perforation of the atrial wall could be seen. The orifice of the coronary sinus was enlarged. The fossa ovalis was 2.5 cm. in diameter and closed. The pectinate muscles of the atrial appendage were hypertrophied. No mural thrombi were seen. The tricuspid valve measured 13.5 cm. in circumference (normal 12 cm.). The tricuspid leaflets showed multiple eccentric vegetations varying in size from 0.7 to 0.2 cm. Some of the

vegetations were located on the valve and its free edge; others on the chordae tendineae (Fig 1). No perforation of the leaflets or rupture of the chordae tendineae were noted. A section from the tricuspid valve and vegetation showed acute bacterial endocarditis. The vegetation contained numerous polymorphonuclear leukocytes, necrotic debris, fibrin, and platelets. The Brown-Brenn stain showed many gram-positive cocci.

The right ventricle was moderately enlarged. The inflow tract measured 10 cm. and the outflow tract measured 12 cm. The endocardium was intact with no hemorrhages or discolored or perforated areas. The wall of the right ventricle was thick, measuring 0.7 cm. in the distal portion of the pulmonary outflow tract. The papillary muscles were hypertrophied and the trabeculae carneae were moderately widened. The pulmonary ring measured 9.5 cm. in circumference (normal 8.5 cm.). The pulmonary cusps were normal. The left atrium was extremely enlarged and was very thin in places. It measured 7.5 cm. in superior-inferior diameter and 15 cm. in circumference. The mitral ring measured 14 cm. in circumference (normal 10 cm.). The leaflets of the mitral valve were minimally fibrotic for this age with some hemodynamic changes at their free edges. There were no adhesions between the leaflets. The major chorda with its 3 branches inserting into the most medial portion of the anterior leaflet was ruptured (Fig 2). This was an old rupture as judged by the thickened rounded ends of the separated segments of the chorda and histologically by fibrosis and hyalinization of the ruptured ends. The apical portion of the posterior papillary muscle from which the chorda arose showed considerable irregular fibrosis with streaked calcification in its tendinous portion. The left ventricle was enlarged. There was no aneurysm. The inflow tract of the left ventricle was 9 cm. and the outflow tract was 10.5 cm. in length. The thickness of the left ventricular wall was 1.8 cm. at its base and 1.3 cm. at the apex. The papillary muscles were hypertrophied. The trabeculae carneae were thick and widened. The endocardium of the left ventricle was normal. No mural thrombus, discolored area, or perforation were noted. The cir-

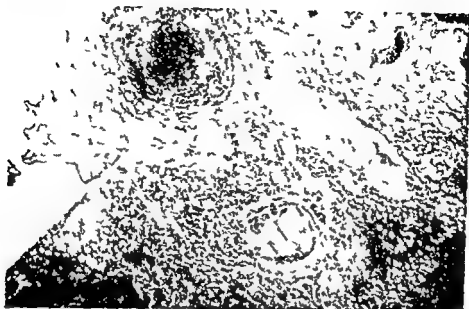


Fig 4 Section through discolored epicardium and myocardium of Fig 3 shows extensive hemorrhage and acute inflammatory process with septic thrombi in the epicardial vessels (Hematoxylin and eosin Original magnification $\times 90$)

cumference of the aortic valve was normal measuring 7.5 cm. The cusps had the usual amount of thickening. Moderate atherosclerosis was noted in the sinuses and around the orifices of the coronary arteries. However, the orifices of the coronary arteries were patent, the right measuring 3 mm and the left 4 mm in diameter. The right coronary artery was dominant forming the posterior descending interventricular branch. All coronary arteries showed moderate atherosclerosis. The lumen of the anterior descending branch was narrowed some 60 per cent in an area 3 cm from the origin of this vessel. The posterior descending branch showed marked narrowing and nearly complete occlusion of its lumen 8 cm from the orifice of the right coronary artery and 7.5 cm from the apex of the heart. A histologic section of the coronary artery from this segment showed marked atherosclerosis, with fresh hemorrhage into the plaque. In the course of this posterior descending branch there was the discolored epicardial area described previously (Fig 3). A section through this area revealed hemorrhagic clotting extending into the midportion of the myocardium. However, the corresponding endocardium was intact. Several sections from this area for histologic study showed an intense acute inflammatory process in the myocardium and epicardium (Fig 4). There were mul-

tiples myocardial abscesses with a polymorphonuclear exudate trailing between muscle fibers and under perivascular tissues. A considerable amount of myocardial fibrosis was present but there was no acute myocardial necrosis indicative of recent infarction. The acute inflammatory process extended into the epicardium where it was associated with hemorrhage. Epicardial arterial branches and veins were crusted up in the inflammatory process with disruption of their walls and in instances with septic luminal thrombi. Brown Brown stains for bacteria showed gram positive cocci intra- and extracellularly in the myocardial and epicardial exudate including the disrupted epicardial blood vessels. Sections from the different portions of the myocardium elsewhere failed to reveal acute inflammatory changes.

The right lung weighed 650 Gm and the left weighed 450 Gm. Their pleural surfaces were normal. The bronchial tree was intact except for some lower lobe branches which were moderately dilated. The pulmonary arteries revealed mild atherosclerotic changes. There were no thrombi. Some dark reddish frothy fluid could be expressed from the parenchyma. There was some peribronchovascular emphysema in the right upper lobe.

The liver weighed 1,640 Gm with a dark reddish color and firmer consistency than

usual Histologically acute and chronic passive congestion were present with minimal central and sublobular perivenous fibrosis

The first portion of the duodenum showed multiple subacute ulcers the largest 1 cm in diameter The second portion of the duodenum had a diverticulum 3.5 cm in width There was no gastrointestinal hemorrhage

Each kidney weighed 150 Gm The surface was moderately granular Histologically there was vacuolization of proximal and distal tubular epithelium However, there were no acute tubular necroses The interstitial tissues in the cortico medullary junctional areas were infiltrated with lymphocytes and polymorphonuclears There were some hyaline and pigmented casts in Henle's loops

One parathyroid showed adenomatous hyperplasia made up largely of chief cells

The brain showed a recent hemorrhagic infarct in the right parietal lobe measuring 2.4 by 2.0 cm In addition microscopically there was an abscess in the mid brain near the *substantia nigra* Gram positive cocci were identified in this microscopic lesion stained by Brown Brenn The pituitary gland was normal grossly but microscopically there were 3 focal areas of infarction with bland appearances No bacteria could be demonstrated in these lesions

DR ARABOWER We felt that the sudden onset of heart failure in 1956 was due to rupture of a chorda tendinea to the medial portion of the anterior leaflet of the mitral valve Such rupture is most frequently associated with bacterial endocarditis less so with rheumatic endocarditis infarct of the papillary muscles Marfan's syndrome or trauma to the chest None of these conditions appears to be the underlying factor in the present case It may be assumed therefore that in the present instance the rupture was spontaneous This is an uncommon occurrence When it does occur however it is in older individuals and the chordae to the posterior leaflet of the mitral valve are the ones that rupture rather than those to the anterior leaflet We are inclined to believe that from 1956 onwards the patient had a considerable degree of mitral insufficiency with left ventricular enlargement and hypertrophy

We found no evidence at the time of death of either a distinctive area of old myocardial infarction or of a ventricular aneurysm There was myocardial fibrosis but it was patchy and particularly in the posteroinferior part of the heart supplied by the distal portion of the posterior descending coronary artery We shall have occasion to refer to this area of myocardium a bit later in this discussion

We would interpret the eventful 3 days following cardiac catheterization as follows Since we were unable to find any evidence for perforation of atrial or aortic walls we believe that the shock which occurred during cardiac catheterization was not due to hemorrhage into the pericardial sac We would ascribe it to some other mechanism We are inclined to believe that the period of shock in one or other way contributed to the renal failure although there was little anatomical evidence at the time of death of acute tubular necrosis Much of this failure may have been due to extrarenal factors However the necroses of the anterior pituitary and the hemorrhagic infarct of the left parietal lobe of the brain could be related to shock and Levophed therapy Aside from shock the major complication following cardiac catheterization was sepsis We would ascribe the source of the sepsis to the central venous catheter In the first place the urinary bladder was clean and while there was slight traumatic hemorrhage due to the Foley catheter there was little evidence for an acute cystitis Second blood cultures taken on 4 consecutive days following catheterization yielded positive cultures for *Staphylococcus aureus* Third the acute bacterial endocarditis involving the tricuspid valve was due to coccal and not gram negative organisms Pyemic lesions were however restricted to one small area in the mid brain and to a larger lesion in the posteroinferior wall of the myocardium In this area of the myocardium there was a diffuse suppurative and hemorrhagic process with abscesses and with readily demonstrable coccal organisms in the lesions There was extension of the suppurative and hemorrhagic process to the overlying epicardium with involvement of the coronary arterial and venous branches In fact some of the vessels contained colonies of organisms We would

ascribe the hemopericardium which we found at the time of death to seepage from these disrupted inflamed epicardial vessels. We are at a loss to explain the localization of the pyemic process to an area of myocardium, the seat of appreciable earlier myocardial scarring, except that the markedly narrowed area of the posterior descending coronary artery demonstrated a fresh hemorrhage into the atherosclerotic plaque. This additional narrowing may have favored bacterial localization by altering the pattern of the blood flow in the distal part of the artery and by the increased ischemic state of that portion of the heart.

The immediate cause of death appeared

to be related in the main to sepsis and the hemopericardium. There was no anatomical evidence for disseminated intravascular thrombosis.

DIAGNOSIS Old spontaneous rupture of chorda tendinea of mitral valve with mitral insufficiency and marked left ventricular hypertrophy and enlargement. Acute staphylococcal bacterial endocarditis of the tricuspid valve ascribable to a central venous catheter with pyemic involvement of a localized area of myocardium and epicardium, leading to disruption of walls of the epicardial vessels with resultant massive hemopericardium.

Fundamentals of clinical cardiology

Mechanisms of cardiac arrhythmias From hypothesis to physiologic fact

Alfred Pick M.D.*
Chicago Ill

In 1959 Drs. Katz and Langendorf and I¹ pleaded for closer cooperation between experimental electrophysiologists and clinical electrocardiographers in order to verify hypothetical concepts and to attack some unresolved problems in interpreting disorders of the rhythm of the human heart. Today we can state that our hopes have been surpassed far beyond our expectations. Indeed the role of the two groups of investigators has been reversed. Clinical applicability of results achieved in ingeniously devised animal experiments is being pointed out to an increasing extent in the writings of electrophysiologists. On the other hand the clinician can now turn to experimentation on the human heart thanks to the development of harmless and precise methods permitting controlled pacing of the heart chambers, the recording of action potentials from the specific conduction system and particularly the combined application of the two techniques.

Among the great number of hypothetical mechanisms that have been tested in this manner twenty that are reproducible in man by cardiac pacing have been dealt with in a previous communication.² For this presentation I selected the following seven:

- 1 Manifestations of concealed conduction
 - 2 Re entry pathways in the atria and/or the A-V junction
 - 3 Concealed re entry in the A-V junction (atypical Wenckebach periods)
 - 4 The question of supernormal conduction
 - 5 Pathways for ventricular pre excitation
 - 6 Dependence of aberrant conduction on cycle length
 - 7 Impulse formation and Mobitz Type I block in the ventricular conduction system
- By correlation of clinical records with their experimental counterparts some observed in our own laboratory but mostly taken from the work of other investigators I shall try to demonstrate how well or how poorly our conjectures have stood up to experimental testing. I shall mention some remaining areas of controversy and shall also include speculations awaiting experimental verification in the future.

Manifestations of concealed conduction

Fig 1(A) is a reproduction of part of a record published by Simon and Langen

Presented at the International Symposium on Reentry Arrhythmias organized by the Department of Cardiology and Clinical Physiology of the University of Amsterdam March 23 and 24 1972 Amsterdam.
Received for publication August 21 1972.
Reprint requests to Dr Alfred Pick Cardiovascular Institute, Michael Reese Hospital and Medical Center, Chicago Ill 60616.
Dr Pick is Professor of Cardiology, Institute of Michael Reese Hospital and Medical Center, Chicago Ill.

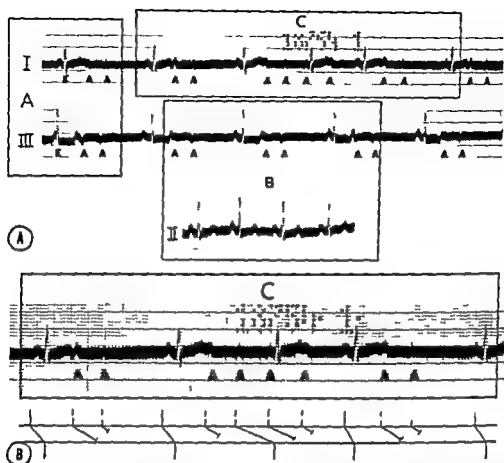


Fig 1. (A) Sinus rhythm with repetitive nonconducted atrial premature ectopic beats. Frames I and B are discussed in text. Frame C (with diagram) is reproduced in (B) showing repetitive nonconduction of atrial premature beats due to concealed A-V conduction. The length of the oblique lines in the diagram indicates the extent of penetration of the A-V junction by ectopic atrial impulses. (Fig 1 I is reproduced from Katz, I. N. and Pick, A. Clinical Electrophysiology, Part I: The Arrhythmias, Philadelphia, 1956, Lea & Febiger. Reproduced with permission.)

dorf³ in 1949. Leads I and III show repetitive nonconducted atrial premature systoles each labelled A while in Lead II the sinus rhythm is undisturbed. Three different points of interest are framed and labelled A, B, and C. Frame A shows a fixed coupling of 0.32 sec of the second premature beat to the first. On this basis a re-entry mechanism in the atria was assumed. This possibility was documented experimentally by Han and associates⁴ who proved that the sinus node was included in the re-entry path. However, this does not seem to apply to our clinical case. In Frame B it can be seen that the sinus interval of 1.66 sec, including the two premature P waves, equals two undisturbed sinus cycles. In other words, that the ectopic atrial impulse is interpolated. That an ectopic atrial impulse capable of producing an atrial echo may or may not enter the sinus node and reset its cycle has been

demonstrated in man experimentally by Goldreyer and Damato.⁵ In our clinical case the entrance block of the premature impulse is bidirectional toward the sinus as well as toward the A-V junction.

However, the integrative block is not always complete as shown in Frame C reproduced with a diagram in Fig 1 (B). In the latter failure of the ventricles to respond to the second of a pair of premature atrial impulses is attributed to concealed penetration of the A-V junction by the first of the pair. Yet this takes place to a varying extent. With less deep penetration of the first premature atrial impulse—in the middle of the record—the subsequent ectopic impulse can reach the ventricles only with great delay.

We were able to reproduce this at a much later time in another patient by atrial pacing as illustrated in Fig 2.⁶ When an extra impulse X was interposed into a

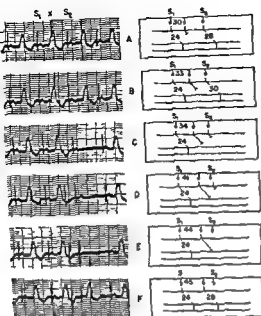


Fig 2 Concealed A-V conduction in paired artificial atrial pacing S_1 Basic driving stimuli X interpolated extrasystoles numbers sec/100 (Fig 2 is reproduced from Langendorf R Pick A Edelstein L and Katz L N Experimental demonstration of concealed A-V conduction in the human heart *Circulation* 32:386 1965 Reproduced by permission of the American Heart Association Inc)

stimulation cycle progressive lengthening of the interval S_1 determined the depth of its concealed conduction and thereby the success or failure of conduction of S_2

In the same case atrial pacing was used to reproduce in man the classical experiment carried out in 1925 by Lewis and Master⁷ on concealed A-V conduction in dogs A comparable result was observed in our experiment in human subjects (Fig 3)⁸ A 2:1 ventricular response with a P-R of 0.28 with rapid atrial pacing changed to a 1:1 response with a shorter P-R of 0.24 when the pacing rate was one half The assumption is that the longer P-R is the aftereffect of concealed conduction into the A-V node of the apparently blocked stimulus We could verify this interpretation by repeating such an experiment with a His bundle recording (Fig 4) In Panel B during rapid pacing of the atrial alternate impulses (S_1) are not followed by a His deflection while the A-H interval of conducted impulses measures 140 msec Thus there is a 2:1 block proximal to H presumably in the A-V node At half the

stimulation rate in Panel C there is a ventricular response to each stimulus with the A-H interval reduced to 110 msec This is similar to that of the control Panel A obtained during sinus rhythm at a rate of 70

Perhaps the most convincing documentation of the concealment of an impulse within the A-V junction was recently provided in the case of so-called pseudo A-V block (Fig 5) A clinical instance the second in our material since the original description by Langendorf and Mehler⁹ 25 years ago shows in Lead II sudden prolongation followed by failure of conduction in a regular sinus rhythm As indicated in the diagram this is attributed to after effects of two premature A-V junctional discharges blocked in both directions This interpretation is supported by Lead III, where the interpolated junctional discharge is conducted to the ventricles with aberration Documentation of such a spontaneously occurring mechanism was provided by Rosen and associates¹⁰ in a clinical case using His bundle recordings and experimentally by Moore and associates¹¹ in the rabbit heart recorded through a microelectrode impaled into the A-V junction

Re entry pathways in atria and/or A-V junction

The combination in man of cardiac pacing and His bundle recordings¹² has made it possible to confirm a number of deductive and empirical interpretations of standard clinical records that have important prognostic and therapeutic implications For example Mobitz Type I A-V block can be distinguished from Type II the location of the block in the conduction system can be ascertained and combinations of block can be recognized^{13,14} Likewise continued reciprocal beating of atria and ventricles has been identified as the cause of many—although not all—supraventricular paroxysmal tachycardias^{15,16} Learning on convincing clinical experimental data^{17,18} we are able to understand the spontaneous initiation and termination of paroxysms in physiologic terms One such instance is illustrated in Fig 6 In the upper strip a junctional circus movement can be postulated to be interrupted when a spontaneous

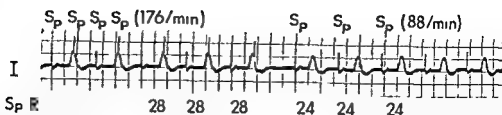


Fig 3 Concealed A V conduction in rapid atrial pacing. *Sp* Atrial response to pacer stimulus (Fig 3 is reproduced from Langendorf R, Pick A, Edelstein A, and Kitz L N. Experimental demonstration of concealed A V conduction in the human heart. *Circulation* 32:386, 1965. Reproduced by permission of the American Heart Association, Inc.)

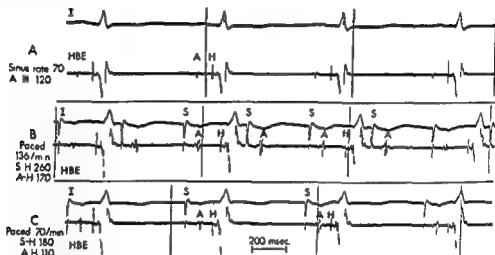


Fig 4 Concealed A V conduction demonstrated by His bundle recording during rapid atrial pacing (B). Panel A Control before pacing. *S* Stimulus artifact in His bundle electrogram (HBE). *A* atrial potential. *H* His potential. *A-H* transnodal conduction time (approximate).

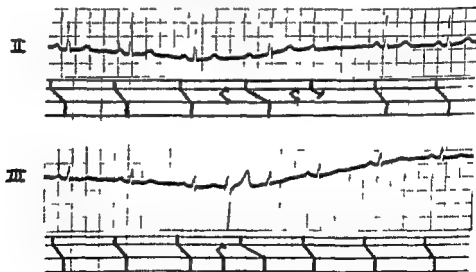


Fig 5 First degree and Type I second degree A V block initiated by blocked A V junctional premature beats

premature atrial discharge at the upper arrow, invades and blocks the circus path; it is resumed in the lower strip when a similar spontaneous atrial ectopic impulse at the lower arrow, conducted with delay to the ventricles, initiates repetitive alternating echos of atria and ventricles. We

also learned why in Type I A V block paroxysms of reciprocating tachycardia prevent the completion of a Wenckebach period once a critically prolonged P-R is reached (Fig 7); it sets the stage for the first as well as consecutive re-entry processes in the A V junction.^{16,27} Another

J J May 86
A mb

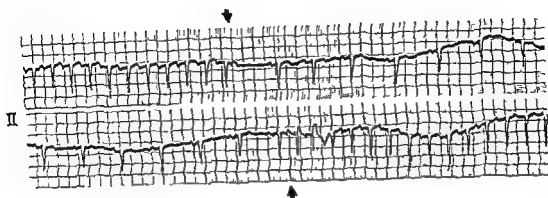


Fig 6 Termination (↓) and resumption (↑) of reciprocating supraventricular paroxysmal tachycardia by atrial premature beats. The right bundle branch block contour of the first and second beat of the recurrent tachycardia is due to aberrant ventricular conduction.

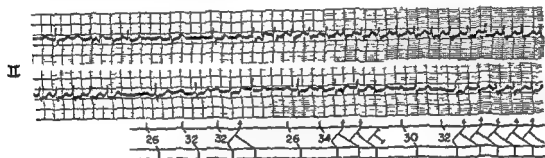


Fig 7 Repetitive reciprocating supraventricular tachycardia initiated by critical (0.32 to 0.34 sec) P-R prolongation in Type I second degree A-V block (caused by digitalis).

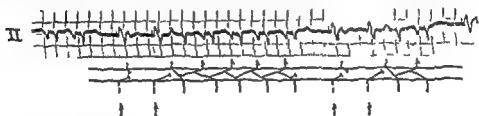


Fig 8 Intermittent reciprocating supraventricular tachycardia initiated by prolonged A-V conduction time of ventricular captures during A-V dissociation resulting from ventricular pacing.

more rare example of the crucial role of slowed A-V conduction in the initiation and perpetuation of atrial and ventricular echoes is shown in Fig 8. During demand pacing of the ventricles indicated by the pairs of upright arrows, A-V dissociation is replaced transiently by atrioventricular reciprocation subsequent to the prolonged P-R interval of a ventricular capture by a sinus impulse.

However, not all known varieties of junctional tachycardia have an experimental counterpart. Continued reciproca-

tion remains to be proved in cases with a bidirectional exit block. Fig 9 shows an example with two diagrams which may or may not require amendment. In a junctional tachycardia there is repetitive group beating of the ventricles and atria corresponding to a 5:4 antegrade and a 3:3 retrograde Type I block. In the left diagram a continuous regular junctional discharge is assumed and double block in retrograde direction attributed to concealed backward penetration of each fourth junctional impulse. In the right diagram

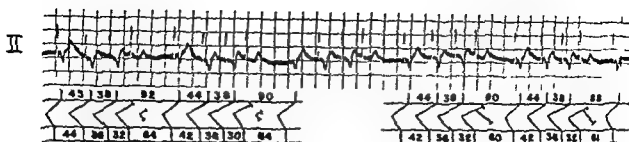


Fig 9 A V junctional tachycardia with Wenckebach periods of forward and retrograde conduction For explanation of the two diagrams see text

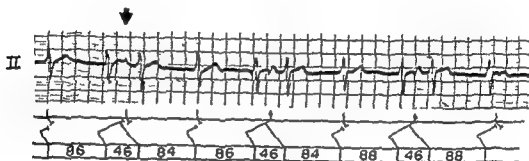


Fig 10 Subatrial re entry in A V junctional rhythm with reciprocal beating

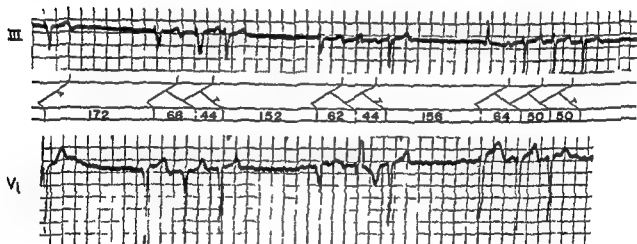


Fig 11 Subatrial re entry in A V junctional rhythm during atrial fibrillation (A V dissociation) in Lead V1

concealed subatrial antegrade re entry is assumed, initiated by the prolonged concealed retrograde conduction which resets the cycle of the junctional focus

Be that as it may such not too rare cases have a bearing on the unsettled dispute^{40, 49} as to whether or not the atria are a necessary link for reciprocal beating of the ventricles or to put it in physiologic terms whether functionally dissociated pathways converge to a common path not only in the distal but also in the proximal A V junction

Other records lead us to believe that in ventricular echoes the reversal of the retrograde impulse takes place at a subatrial level For instance in Fig 10 showing a case of reciprocal beating at the arrow a sinus P wave is sandwiched between the first couple in contrast to clearly retrograde P waves in the second and third couple Perhaps less convincing but plausible is our interpretation of Fig 11 Similar groups of two or three fast beats follow junctional escapes In Lead III the retrograde P waves identify the arrhythmia as reciprocal

1 me E6
N mbe

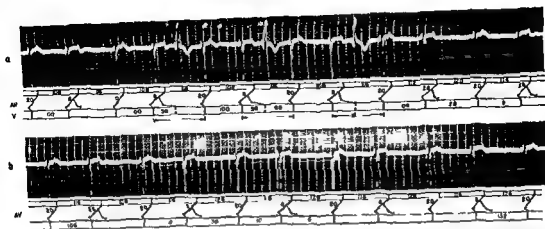


Fig 12 Concealed (blocked) reciprocal beating in A-V junctional rhythm with delayed retrograde conduction. Note that in *b* the longer R-P interval of 0.28 s is always followed by prolongation of the manifest junctional cycle from 1.08 to 1.36 sec

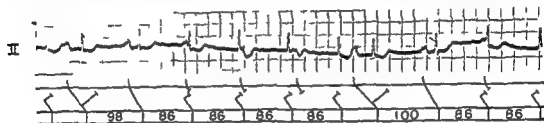


Fig 13 Concealed retrograde re-entry (attempt at atrial echo) in incomplete A-V dissociation which developed after acceleration of A-V junctional impulse formation by digitalis

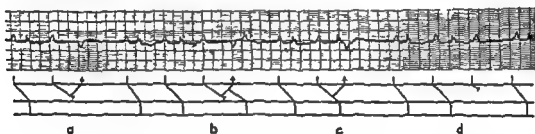


Fig 14 Second-degree A-V block (Type I) with concealed A-V conduction with three atrial echoes and one ventricular echo

beating. In Lead V_1 on the next day when atrial fibrillation had developed and the atria became unresponsive the assumed reciprocal path would have to be below the atronodal junction.¹⁹

Concealed re-entry in A-V junction

Reciprocal beating is one of the manifestations in the human heart of the experimentally proved existence of at least two A-V transmission systems having different functional properties.^{17, 18} These potential re-entry pathways may not be utilized to

their entire length and the circus movement may be incomplete. Thus in 1950²⁴ we explained a curious case (Fig 12) of slow junctional rhythm in which R-P intervals and junctional cycles alternated concomitantly by concealed antegrade re-entry postponing the next junctional discharge. The clue was found in the upper record of another day which showed that the prolongation of the ventricular interval to 1.24 was the result of junctional re-entry actually completed in the middle but concealed at both ends of the record. By the

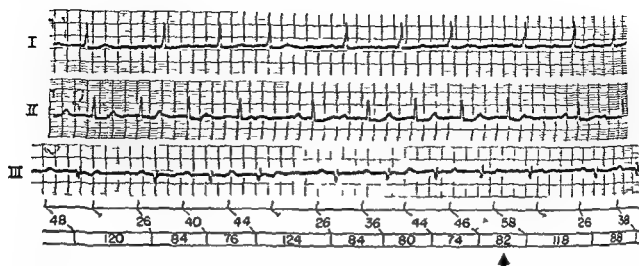


Fig 15 Concealed re entry in the AV junction (attempt at atrial echo) causing an atypical Wenckebach period

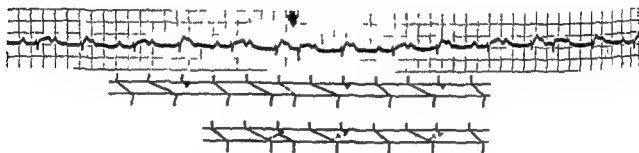


Fig 16 Atypical Wenckebach periods due to a double AV path (upper diagram) or concealed retrograde conduction followed by manifest anterograde re entry (lower diagram) in the AV junction

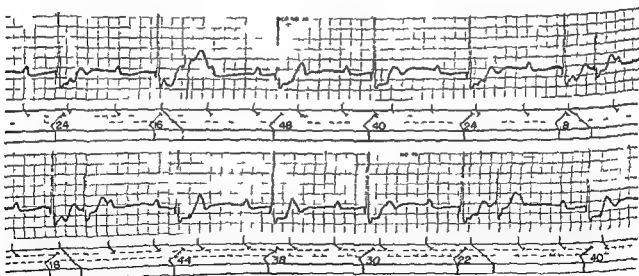


Fig 17 Incomplete AV dissociation due to advanced unidirectional AV block. Concealed retrograde conduction of idioventricular impulses permits ventricular capture by sinus impulses (with aberrant ventricular conduction) only during a supernormal pause of AV conduction

same token in another case (Fig 13) with incomplete AV dissociation we explained the failure of a junctional escape to occur at the expected time of 86 by a concealed retrograde re entry attempt induced by

prolonged integrate conduction of early ventricular captures.⁴⁶ Experimental data now available in the literature^{45, 47} allow us to interpret with confidence other variants of such mechanisms. — example,

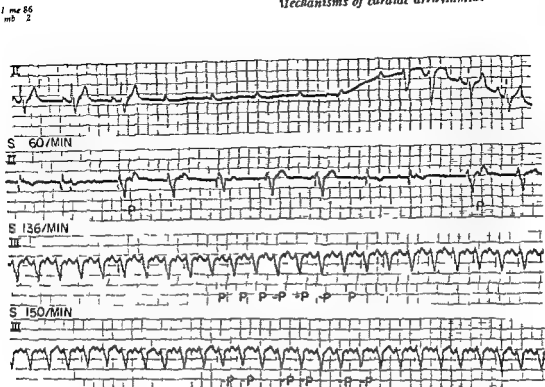


Fig 18 Undirectional A V block in Stokes Adams syndrome. The first strip shows a period of prolonged ventricular asystole followed by two ectopic ventricular beats before A V conduction is resumed. Note constancy of P R in conducted beats (Type II antegrade A V block). In the second strip such intermittent periods of asystole are prevented by responses to transvenous ventricular asynchronous pacing at a rate (S) of 60. Pacer impulses failing at long distances after a P wave were conducted backwards to the atria (P). When the pacing rate was increased to 136 (third strip) each artificial impulse reached the atria at a constant R P interval of 0.24 sec. Thus the A V block was proved to be undirectional. With further acceleration of the pacing rate to 150 the atria respond at a 3:2 ratio with a Wenckebach phenomenon, the second R P lengthening to 0.26 sec. This region of Type I retrograde block was probably proximal to the Type II antegrade block.

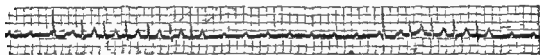


Fig 19 Sudden protracted ventricular arrest in Type II second degree A V block. A V conduction is resumed following spontaneous escape of a subsidiary ectopic pacemaker.

some atypical features of atrioventricular Wenckebach periods.⁴⁹

An unusual case is illustrated in Fig 14 in which four different events are labelled *a* to *d* for presentation. Part *d* shows a simple 3:2 Wenckebach period. In *a* to *c* the sequence of regular sinus impulses is disturbed by atrial echoes induced by greater delay in antegrade conduction. These echoes occur regardless of whether the initiating sinus impulse reaches the ventricles as in *c* or fails to do so as in *a* and *b*. However, when it fails, the unused portion of the antegrade path provides a potential avenue for a ventricular echo: a sweep that is actually completed in portion *b*.

In the Wenckebach periods shown in Fig 15 the arrow indicates that against the rule the increment of P R is largest before the dropped beat. Previously⁴⁹ we tried to explain this observation by assuming a critical state of A V conductivity before its total failure. Today we can attribute it to the aftereffect of an attempted reentry via a retrograde path that has opened up after sufficient delay of antegrade conduction is indicated in the diagram. Fig 16 illustrates an opposite event. In the middle of a Wenckebach period at the arrow the P R unexpectedly shortens before the period is resumed and completed. Originally⁴⁹ we thought that this might be

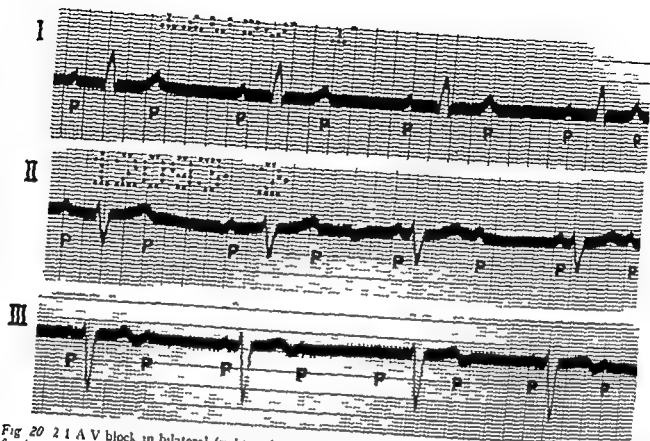


Fig 20 2:1 A-V block in bilateral (right and left anterior fascicular) bundle branch system block. Note two fixed ranges (0.16 and 0.34 sec) of P-R intervals in the conducted beats

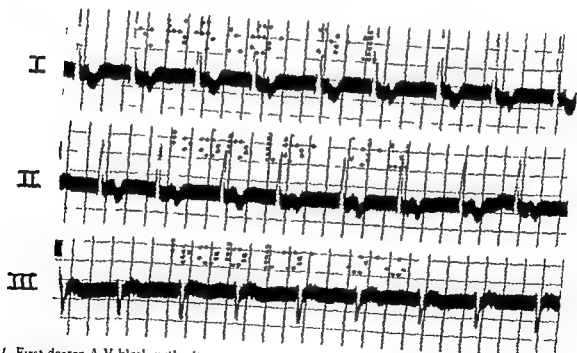


Fig 21 First degree A-V block with alternation of P-R intervals (causing ventricular bigeminy). For interpretation see text and Fig 22

due to supernormal conduction through depressed A-V junctional tissues. In the light of advanced knowledge other interpretations appear more likely. In the upper diagram two longitudinally dissociated conduction pathways with different con-

duction speeds are considered. The lower diagram indicates that the premature QRS is unrelated to the P wave preceding it. Rather it is a ventricular echo reflected from an abortive (concealed) attempt at an atrial echo. If true this interpretation

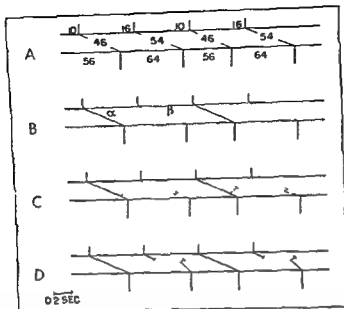


Fig 22 Diagrammatic interpretation of Fig 21. The numbers in diagram A represent (in sec./100) R P intervals in the upper level, P R intervals in the middle level, and R R intervals in the lowest level. SVP, supernormal phase of A V conduction. Two A V pathways (α and β) are indicated in diagrams B and C by solid and broken diagonal lines. B, Double path 2:1 blocked, α shorter RP, faster conduction; C, Double path 2:1 blocked, α shorter RP, slower conduction; D, 2:1 block with junctional escape.

serves as evidence against inclusion of the atria into the re-entry path in reciprocal beating (see above).

Supernormal conduction

The concept of supernormal A V conduction is hypothetical and indeed was severely criticized by Moe and his associates.⁴¹ It is unlikely in the case of unexpected shortening of conduction time of successive impulses traveling in the same direction.^{42,43} However, we still consider the mechanism of supernormal conduction in advanced A V block (Fig 17) when captures occur exclusively and predictably during a narrow, limited period early in the ventricular cycle. In the example shown, the range for A V conduction expressed in R P intervals is 0.16 to 0.22 sec. We attribute this to backward penetration of ventricular impulses transiently converting a region of unidirectional conduction into one of supernormal conduction.⁴⁴ We were able to reproduce this at will in a number of cases of advanced A V block requiring ventricular pacing during slow pacing rates and to prove the existence of unidirectional block by driving the ventricles faster than the spontaneous rate of the sinus node (Fig

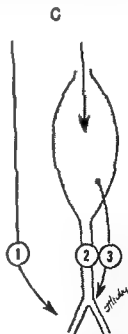
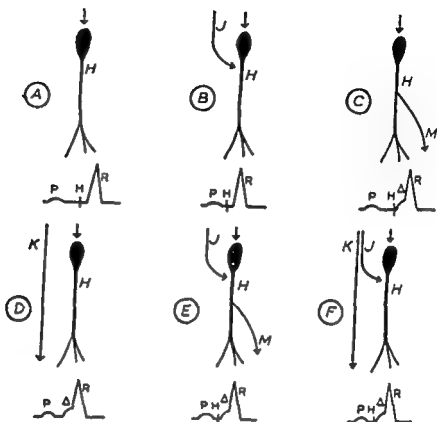


Fig 3 Schematic representation of pathways for A V conduction in ventricular pre-excitation. 1, Total A V bypass; 2, normal path through A V node and bundle of His; 3, partial A V bypass (Fig 23 is reproduced from Katz, L. N. and Pick, A. Clinical Electrocardiography, Part I, The Arrhythmias, Philadelphia, 1956, Lea & Febiger. Reproduced with permission).



Diverses combinaisons possibles des voies de pré-excitation

- a) Conduction auriculo ventriculaire normale
 b) Syndrome du P R court sans onde delta (Fibres de James)
 c) Syndrome de WPW avec espace P R normal (Fibres de Mahaim)
 d) Syndrome de WPW avec P R court et espace P H normal par voie de pré excitation auriculo ventriculaire directe (Faisceau de Kent)
 e) Syndrome de WPW avec P R court et P H court, par association de fibres de James et de fibres de Mahaim
 f) Syndrome de WPW avec P R court et P H court par association d'un faisceau de Kent et de fibres de James
 (Abreviations H = faisceau de His J = fibres de James M = fibres de Mahaim K = faisceau de Kent)
 Pour la discussion voir le texte

Fig 27 A schematic correlation of the manifestations in the electrocardiogram of the function of various total and partial A V bypasses J James fibers K Kent fibers M Mahaim fibers H His bundle potential and its temporal relation to P and R waves with normal A V conduction (A) and in variants of ventricular pre excitation (B F) see text (Fig 1 in Coumel and associates⁶⁶ reproduced with permission from Coumel Wayneberger Slama and Bouvrain Acta Cardiol 26 188 1971)

bundle branch block (Fig 28) we attributed persistence of the latter to a bypass pre exciting the contralateral left ventricle⁶⁷ But this assumption was untenable in another case (Fig 29) of intermittent right bundle branch block associated with pre excitation of Type B In the framed Lead V₁ a deep negative delta wave is followed by a large late R wave Here right sided pre excitation did not preclude the bundle branch block in the ipsilateral ventricle Pending a better explanation based on anatomic and physiologic study of such cases, we propose that a complete right sided A V bypass could insert proximal to

the region of block in the right ventricular conduction system

Dependence of aberrant conduction on cycle length

Turning now to some other problems of abnormal intraventricular conduction we will consider causes of aberrant conduction of escape beats An example is shown in Fig 30 In a digitalized patient with atrial fibrillation regular escape beats the smaller QRS complexes in each lead show a higher degree of right bundle branch block than the ones conducted faster The

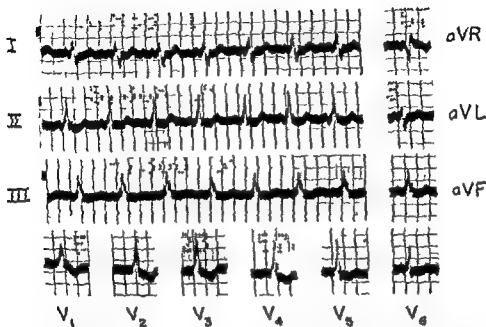


Fig 28 Ventricular pre-excitation Type A in the presence of a right bundle branch block

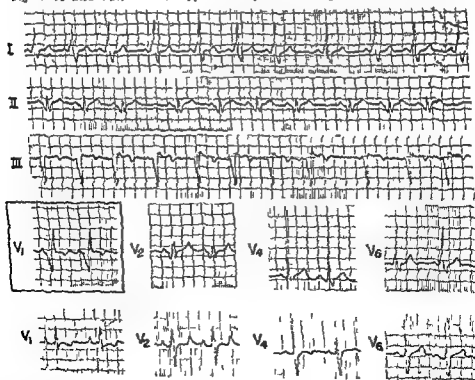
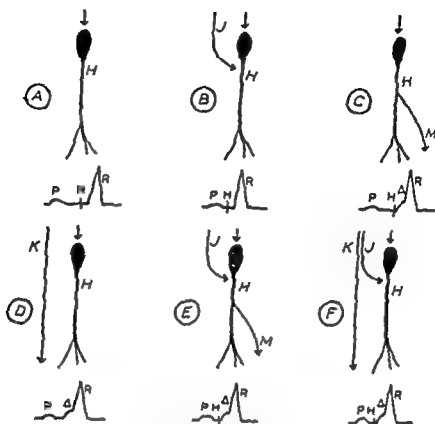


Fig 29 Intermittent ventricular pre-excitation Type B in the presence of right bundle branch block. Pre-excitation is seen in the limb leads and upper series of chest leads and has disappeared in the lower series of Leads V1 to V6.



- Diverses combinaisons possibles des voies de pré-excitation
- Conduction auriculo-ventriculaire normale
 - Syndrôme du P-R court sans onde delta (Fibres de James)
 - Syndrôme de W-P-W avec espace P-R normal (Fibres de Mahaim)
 - Syndrôme de W-P-W avec P-R court et espace P-H normal par voie de pré-excitation auriculo-ventriculaire directe (Faisceau de Kent)
 - Syndrôme de W-P-W avec P-R court et P-H court, par association de fibres de James et de fibres de Mahaim
 - Syndrôme de W-P-W avec P-R court et P-H court par association d'un faisceau de Kent et de fibres de James
- (Abréviations: H = faisceau de His, J = fibres de James, M = fibres de Mahaim, K = faisceau de Kent)
- Pour la discussion voir le texte

Fig 27 A schematic correlation of the manifestations in the electrocardiogram of the function of various total and partial A-V bypasses: J James fibers, K Kent fibers, M Mahaim fibers, H His bundle potential and its temporal relation to P and R waves with normal A-V conduction (A) and in variants of ventricular pre-excitation (B-F) see text (Fig 1 in Coumel and associates⁴⁸ reproduced with permission from Coumel, Waynberger, Slama and Bouvraie, *Acta Cardiol* 26:188, 1971)

bundle branch block (Fig 28) we attributed persistence of the latter to a bypass pre-exciting the contralateral left ventricle.⁴⁷ But this assumption was untenable in another case (Fig 29) of intermittent right bundle branch block associated with pre-excitation of Type B. In the framed Lead V₁ a deep negative delta wave is followed by a large late R wave. Here right sided pre-excitation did not preclude the bundle branch block in the ipsilateral ventricle. Pending a better explanation based on anatomic and physiologic study of such cases, we propose that a complete right sided A-V bypass could insert proximal to

the region of block in the right ventricular conduction system.

Dependence of aberrant conduction on cycle length

Turning now to some other problems of abnormal intraventricular conduction we will consider causes of aberrant conduction of escape beats. An example is shown in Fig 30. In a digitalized patient with atrial fibrillation regular escape beats the smaller QRS complexes in each lead show a higher degree of right bundle branch block than the ones conducted faster. The

V 1 me 86
A mb 2

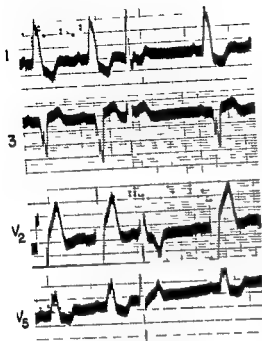


Fig 33 Left bundle branch block (QRS 0.16 sec) with premature beats of bilateral (right and left anterior fascicular) bundle branch block configuration having less QRS prolongation (0.14 sec)

six explanations that have been offered for this apparent paradox are listed in Table I. The first and the last three of the listed mechanisms are purely hypothetical and may apply only exceptionally. The effects of phase 4 depolarization have been determined by Singer and co workers⁷⁶ who have performed microelectrode studies of automatic Purkinje cells. Aberration of RST depends on the magnitude of the transmembrane potential at the time of an ectopic impulse. This take-off potential may be reduced as a result of (1) incomplete repolarization in the case of a premature impulse or (2) diastolic phase 4 depolarization with a late impulse. The latter mechanism is most likely in cases such as that illustrated in Fig 31. Normal intraventricular conduction is maintained up to a cycle length of 112 is slightly aberrant with a cycle of 116 and totally aberrant with cycles equal to or longer than 120. In other cases Rosenbaum's⁷¹ concept of fascicular origin of ectopic beats can be more readily applied. In a case of atrial fibrillation (Fig 32) escape beats can be

identified in the routine electrocardiogram on top by their constant long cycle, as well as by the pattern of a posterior fascicular block. Their ventricular origin presumably in the left anterior fascicle could be documented by the His electrogram. They lack an H deflection that precedes the conducted beats and the fusion beat (F) at a normal H-V interval (lowest panel).

However the possibility of supernormal conduction in depressed bundle branches cannot be entirely dismissed. One pertinent example is shown in Fig 33. In a case of left bundle branch block premature beats have a typical contour of a bifascicular right and left anterior conduction defect. Conceivably the ectopic beats could arise in the posterior fascicle of the left bundle branch system distal to a region of high predivisional unidirectional block. However it could also be postulated that supernormal conduction through the main left bundle unmasks a latent block in the right bundle branch system and left anterior fascicle.⁷⁷

Impulse formation and Mobitz Type I block in ventricular conduction system

One more aspect of abnormal intraventricular conduction will highlight how new techniques have borne out hypotheses (Fig 34). In a case of intermittent ventricular bigeminy after progressive lengthening of the coupling interval of ventricular premature beats from 54 to 70 we speculated on an intraventricular Wenckebach mechanism⁷⁸ located either in a re-entry path or at the Purkinje myocardial junction. Support for this assumption was found in a patient with pacemaker dysfunction (Fig 35). Progressively longer stimulus to response intervals preceded a nonresponse. The prolonged latency after two successive failures as seen at the end of the record could be attributed to decremental propagation of the pacer stimulus.

That a Wenckebach type of block can indeed occur between contractile myocardium and Purkinje fibers was demonstrated in 1961 by Alanis and associates⁷⁹ in Mexico in the rapidly driven dog heart. Recently Greenspan and associates⁸⁰ demonstrated this in hypoxic canine Purkinje fibers as well as in excised human ventricular tissue.

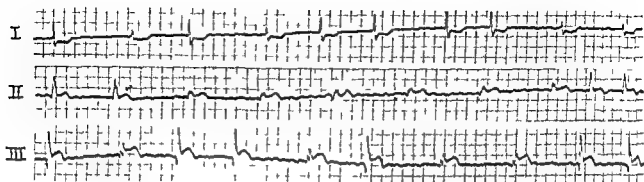


Fig 30 Atrial fibrillation with slow ventricular response and A V junctional escapes with aberrant ventricular conduction (smaller QRS complexes in each lead) causing temporary A V dissociation in the middle of Lead II

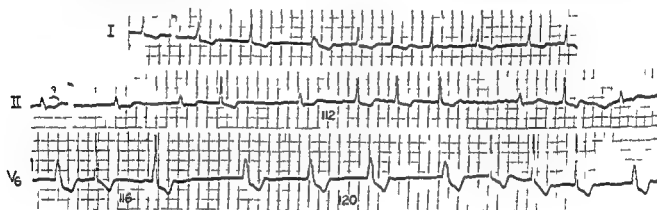


Fig 31 Intermittent (bradycardia dependent) left bundle branch block in atrial fibrillation and left ventricular hypertrophy The degree of the intraventricular conduction disturbance is determined by the cycle length

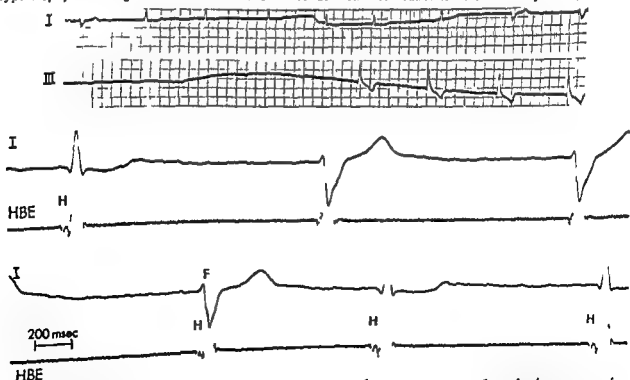


Fig 32 Atrial fibrillation with a slow ventricular response and repetitive escape of a subsidiary pacemaker probably located in the anterior fascicle of the left bundle branch His bundle recording (HBE) reveals that H potentials precede only conducted beats and a fusion beat (F)

- arrhythmia due to multiple premature systoles and concealed A-V conduction without evidence of organic heart disease *Acta Cardiol* 4:54 1949
- 4 Han J, Malozzi A M and Moe G K Sinu-atrial reciprocation in the isolated rabbit heart *Circ Res* 22:335 1968
- 5 Goldreyer B N and Damato A N Sinu-atrial node entrance block *Circulation* 44:789 1971
- 6 Pick A and Langendorf R After effect of paired cardiac excitation in clinical electrocardiograms *Bull N Y Acad Med* 41:606 1965
- 7 Lewis T and Master A M Observations upon conduction in the mammalian heart *Heart* 12:209 1925
- 8 Langendorf R, Pick A, Edelst A and Katz L N Experimental demonstration of concealed A-V conduction in the human heart *Circulation* 39:386 1969
- 9 Langendorf R and Mehlman J B Blocked (nonconducted) A-V nodal premature systoles imitating first and second degree A-V block *Am Heart J* 34:300 1947
- 10 Rosen K M, Rahimtoola S H and Gunnar R M Pseudo A-V block secondary to premature nonpropagated His bundle depolarizations Documentation by His bundle electrocardiography *Circulation* 42:367 1970
- 11 Moore E N, Knoebel S B and Spear J F Concealed conduction *Am J Cardiol* 28:406 1971
- 12 Damato A N, Lau S H, Patton R D, Steiner C and Berkowitz W D A study of atrioventricular conduction in man using premature atrial stimulation and His bundle recordings *Circulation* 40:61 1969
- 13 Wit A L, Weiss M B, Berkowitz W D, Rosen K M, Steiner C and Damato A N Patterns of atrioventricular conduction in the human heart *Circ Res* 27:345 1970
- 14 Wit A L, Damato A N, Weiss M B and Steiner C Phenomenon of the gap in atrioventricular conduction in the human heart *Circ Res* 27:689 1970
- 15 Watanabe Y and Dreifus L S Second degree atrioventricular block *Cardiovasc Res* 1:150 1967
- 16 Damato A N, Lau S H, Helfant R, Stein E, Patton R M, Scherlag H J and Berkowitz W D A study of heart block using His bundle recordings *Circulation* 39:297 1969
- 17 Massumi R A and Ali N Determination of the site of impaired conduction in atrioventricular block *J Electrocardiol* 3:193 1970
- 18 Schuilenburg R M and Durrer D Observations on atrioventricular conduction in patients with bundle branch block *Circulation* 41:967 1970
- 19 Rosen K M, Loeb H S, Chuganua R, Sinno M Z, Rahimtoola S H and Gunnar R M Site of heart block in acute myocardial infarction *Circulation* 42:925 1970
- 20 Narula H S, Scherlag B J, Samet P and Javier R P Atrioventricular block Localization and classification by His bundle recordings *Am J Med* 50:146 1971
- 21 Puech P, Groileau Raoux R, Latour H, Cabasson J, Robin J M, Baisieux C and Gilbert M Diagnostic des blocs tronculaires hisiens par l'enregistrement endocavitare du faisceau de His *Arch Mal Coeur* 65:315 1971
- 22 Haft J I, Weinstock M and DeGuia R Electrophysiologic studies in Mobitz II second degree heart block *Am J Cardiol* 27:682 1971
- 23 Langendorf R Terminology and classification of disturbances of A-V conduction *Bull N Y Acad Med* 47:877 1971
- 24 Rosen K M, Mehta A, Rahimtoola S and Miller H A Sites of congenital and surgical heart block as defined by His bundle electrocardiography *Circulation* 44:833 1971
- 25 Mandel W J, Lozano J, Carascos H and Hayakawa H Coexisting intra and subnodal block An unusual abnormality of atrioventricular conduction *Am Heart J* 82:586 1971
- 26 Schuilenburg R M and Durrer D Conduction disturbances located within the His bundle *Circulation* 45:612 1972
- 27 Hustin A D Mechanisms determining reciprocal rhythm initiated by ventricular premature systoles Multiple pathways of conduction *Am J Cardiol* 31:65 1973
- 28 Moe G K and Mendez C The physiologic basis of reciprocal rhythm *Progr Cardiovasc Dis* 8:461 1966
- 29 Mendez C and Moe G K Demonstration of a dual A-V nodal conduction system in the isolated rabbit heart *Circ Res* 19:378 1966
- 30 Schuilenburg R M and Durrer D Atrial echo beats in the human heart elicited by induced atrial premature beats *Circulation* 37:680 1968
- 31 Schuilenburg R M and Durrer D Ventricular echo beats in the human heart elicited by induced ventricular premature beats *Circulation* 40:337 1969
- 32 Coumel P, Cabrol C, Fabiato A, Gourgon R and Slama S Tachycardie permanente par rythme reciproque I Preuves de diagnostic par stimulation auriculaire et ventriculaire II Traitement par l'implantation intracorporelle d'un stimulateur cardiaque avec entrainement simultané de l'oreillette et du ventricule *Arch Mal Coeur* 60:1830 1967
- 33 Zeft H J and McGowan R L Termination of paroxysmal junctional tachycardia by right ventricular stimulation *Circulation* 40:919 1969
- 34 Janse M J, Van Capell F J L, Freud G E and Durrer D Circus movement within the A-V node as a basis for supraventricular tachycardia as shown by multiple microelectrode recording in the isolated rabbit heart *Circ Res* 28:403 1971
- 35 Wellens H J J Electrical stimulation of the heart in the study and treatment of tachycardias Baltimore 1971 University Park Press
- 36 Goldreyer B N and Damato A N The essential role of atrioventricular conduction delay in the initiation of paroxysmal supraventricular tachycardia *Circulation* 43:697 1971

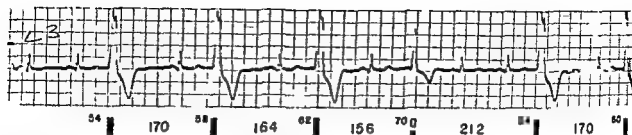


Fig 34 Intermittent ventricular bigeminy. Progressive prolongation of coupling interval (upper small numbers) of ventricular premature beats. The longest coupling (0.70 sec) before intermission approaches the length of the sinus cycle resulting in a ventricular fusion beat (interrupted white/black bar in the diagram). Note that the irregular interectopic intervals (lower larger numbers) have the structure of a Wenckebach period.



Fig 35 Type I exit block of impulses of a failing artificial ventricular pacemaker. In the diagram the dots indicate pacer discharge. Horizontal lines indicate the duration of latency of ventricular responses (↓). Interrupted horizontal lines indicate decremental (concealed) conduction from point of stimulation into surrounding myocardium. (Courtesy of Dr Sidney Goldstein, Rochester, N.Y.)

Table I Potential mechanisms of aberration of escape beats

- 1 Preferential A-V path (origin in Mahaim fibers)^{64, 65}
- 2 Phase 4 depolarization⁷⁰
- 3 Ventricular (fascicular) origin⁷¹
- 4 Functional dissociation within a bundle branch⁷²
- 5 Wedensky facilitation⁷³
- 6 Supernormality of intraventricular conduction^{74, 75}

perfused by potassium in a high concentration

Conclusion and summary

A correlation of deductive interpretations of various disorders of the cardiac rhythm encountered in clinical electrocardiography with mechanisms established in animal and man by the use of modern techniques of registration confirms many but not all previously made assumptions. The present correlation deals with only certain selected mechanisms amenable to experimental testing at the time of preparation of the presentation, in the spring of

1972. In the meantime rapid progress in electrophysiologic techniques has enlarged the possibilities of correlation of clinical presumptions with physiologically established facts. Thus the impact of detection of 'gating' mechanisms operating within the atrioventricular and intraventricular conduction system⁸¹ and the role of slow conduction studied in a new preparation in the genesis of ventricular arrhythmias caused by reentry^{82, 83} and/or parasystolic activity—with and without exit block^{84, 85}—will be included in a future more extensive review article.

The author wishes to thank Drs D. Durrer and H. J. J. Wellens in Amsterdam and Dr P. Coumel and associates in Paris for their kind permission to reproduce Figs. 26 and 27 and Dr S. Goldstein in Rochester, N.Y., for use of his observation in Fig. 35.

REFERENCES

- 1 Pick A, Langendorf R and Katz L N. New aspects and problems in interpreting arrhythmias. Editorial. *Circulation* 20:997, 1959.
- 2 Langendorf R and Pick A. Artificial pacing of the human heart. Its contribution to the understanding of arrhythmias. *Am J Cardiol* 28:516, 1971.
- 3 Simon A J and Langendorf R. complex

- arrhythmia due to multiple premature systoles and concealed A V conduction without evidence of organic heart disease *Acta Cardiol* 4:54 1949
- 4 Han J Malozzi A M and Moe G K Sino-atrial reciprocation in the isolated rabbit heart *Circ Res* 22:355 1968
- 5 Goldreyer B N and Damato A N Sino-atrial node entrance block *Circulation* 44 789 1971
- 6 Pick A and Langendorf R After effect of paired cardiac excitation in clinical electrocardiograms *Bull N Y Acad Med* 41:606 1965
- 7 Lewis T and Master A M Observations upon conduction in the mammalian heart *Heart* 12 209 1925
- 8 Langendorf R Pick A Edelstein A and Katz L N Experimental demonstration of concealed A V conduction in the human heart *Circulation* 32:386 1965
- 9 Langendorf R and Mehlman J B Blocked (nonconducted) A V nodal premature systoles imitating first and second-degree A V block *Am HEART J* 500 1947
- 10 Rosen K M Rahimtoola S H and Gunnar R M Pseudo A V block secondary to premature nonpropagated His bundle depolarizations Documentation by His bundle electrocardiography *Circulation* 42:367 1970
- 11 Moore E N Knoebel S B and Spear J F Concealed conduction *Am J Cardiol* 28 406 1971
- 12 Damato A N Lau S H Patton R D Steiner C and Berkowitz W D A study of atrioventricular conduction in man using premature atrial stimulation and His bundle recordings *Circulation* 40 61 1969
- 13 Wit A L Weiss M B Berkowitz W D Rosen K M Steiner C and Damato A N Patterns of atrioventricular conduction in the human heart *Circ Res* 27:345 1970
- 14 Wit A L Damato A N Weiss M B and Steiner C Phenomenon of the gap in atrioventricular conduction in the human heart *Circ Res* 27 679 1970
- 15 Watanabe Y and Dreifus L B Second degree atrioventricular block *Cardiovasc Res* 1:150 1967
- 16 Damato A N Lau S H Helfant R Stein E Patton R D Scherlag B J and Berkowitz W D A study of heart block using His bundle recordings *Circulation* 39 297 1969
- 17 Massumi R A and Ali N Determination of the site of impaired conduction in atrioventricular block *J Electrocardiol* 3:193 1970
- 18 Schulenburg R M and Durrer D Observations on atrioventricular conduction in patients with bilateral bundle branch block *Circulation* 41:967 1970
- 19 Rosen K M Loeb H S Chuganma R Sinno M E Rahimtoola S H and Gunnar R M Site of heart block in acute myocardial infarction *Circulation* 42 925 1970
- 20 Narula D S Scherlag B J Samet P and Javier R P Atrioventricular block. Localization and classification by His bundle recordings *Am J Med* 50:146 1971
- 21 Puech P Grolleau Raoux R Latour H Cabasson J Robin J M Baisus C and Gilbert M Diagnostic des blocs tronculaires hisiens par l'enregistrement endocavitare du faisceau de His *Arch Mal Coeur* 66:315 1971
- 22 Haft J I Weinstock M and DeGuia R Electrophysiologic studies in Mobitz II second degree heart block *Am J Cardiol* 27 682 1971
- 23 Langendorf R Terminology and classification of disturbances of A V conduction *Bull N Y Acad Med* 47 877 1971
- 24 Rosen K M Mehta A Rahimtoola S and Miller R A Sites of congenital and surgical heart block as defined by His bundle electrocardiography *Circulation* 44 833 1971
- 25 Mandel W J Lozano J Carascos H and Hayakawa H Coexisting intra and subnodal block An unusual abnormality of atrioventricular conduction *AM HEART J* 82:586 1971
- 26 Schulenburg R M and Durrer D Conduction disturbances located within the His bundle *Circulation* 40 612 1972
- 27 Kistner A D Mechanisms determining reciprocal rhythm initiated by ventricular premature systoles Multiple pathways of conduction *Am J Cardiol* 3 365 1959
- 28 Moe G K and Mendez C The physiologic basis of reciprocal rhythm *Progr Cardiovasc Dis* 8 461 1966
- 29 Mendez C and Moe G K Demonstration of a dual A V nodal conduction system in the isolated rabbit heart *Circ Res* 19 378 1966
- 30 Schulenburg R M and Durrer D Atrial echo beats in the human heart elicited by induced atrial premature beats *Circulation* 37 680 1968
- 31 Schulenburg R M and Durrer D Ventricular echo beats in the human heart elicited by induced ventricular premature beats *Circulation* 40:337 1969
- 32 Coumel P Cabrol C Fabiato A Gourgon R and Slama S Tachycardie permanente par rythme réproque I Preuves de diagnostic par stimulation auriculaire et ventriculaire II Traitement par implantation intracorporelle d'un stimulateur cardiaque avec entraînement simultané de l'oreillette et du ventricule *Arch Mal Coeur* 60 1830 1967
- 33 Zeff H J and McGowan R L Termination of paroxysmal junctional tachycardia by right ventricular stimulation *Circulation* 40 919 1969
- 34 Janse M J Van Capell F J L Freud G E and Durrer D Circus movement within the A V node as a basis for supraventricular tachycardia as shown by multiple microelectrode recording in the isolated rabbit heart *Circ Res* 28:403 1971
- 35 Wellens H J J Electrical stimulation of the heart in the study and treatment of tachycardias Baltimore 1971 University Park Press
- 36 Goldreyer B N and Damato A N The essential role of atrioventricular conduction delay in the initiation of paroxysmal supraventricular tachycardia *Circulation* 43 697 1971

- 37 Mandel W J Kermatier A J Blum R L and Hayakawa H Critical prolongation of A V conduction time is the inciting mechanism in reentrant tachycardia J Electrocardiol 11:39 1972
- 38 Goldreyer B N Weiss M B and Damato A N Supraventricular tachycardia initiated by sinus beats Circulation 44:820 1971
- 39 Pick A Langendorf R and Katz L N A V nodal tachycardia with block Circulation 24:12 1961
- 40 Mendez C Han J Garcia de Jalón P D and Moe G K Some characteristics of ventricular echoes Circ Res 13:57 1956
- 41 Mignone R J and Wallace A G Ventricular echoes Evidence for dissociation of conduction and reentry within the A V node Circ Res 19:638 1966
- 42 Goldreyer B N and Bigger J T Jr Site of reentry in paroxysmal supraventricular tachycardia in man Circulation 43:15 1971
- 43 Schuilenburg R M and Durrer D Further observations on the ventricular echo phenomenon elicited in the human heart Is the atrium part of the echo pathway? Circulation 45:629 1972
- 44 Schimroth L Case report Reciprocal rhythm of ventricular origin during atrial fibrillation with complete A V block Br Heart J 32:564 1970
- 45 Pick A and Langendorf R A case of reciprocal beating with evidence of repetitive and blocked reentry of the cardiac impulse Am Heart J 40:113 1950
- 46 Pick A and Langendorf R Recent advances in the differential diagnosis of A V junctional arrhythmias Am Heart J 76:553 1968
- 47 Damato A N Virguese P J Lau S H Gallagher J J and Bobb G Y Manifest and concealed re-entry A mechanism of A V nodal Wenckebach phenomenon Circ Res 30:283 1972
- 48 Castillo C Orlando M and Castellanos A Jr His bundle recordings in atypical A V nodal Wenckebach block during cardiac pacing Am J Cardiol 27:570 1971
- 49 Katz L N and Pick A Clinical electrocardiography Part I The arrhythmias Philadelphia 1956 Lea & Febiger Publishers
- 50 Pick A Langendorf R and Katz L N The supernormal phase of atrioventricular conduction I Fundamental mechanisms Circulation 24:388 1962
- 51 Moe G K Childers R W and Mendeth J An appraisal of supernormal A V conduction Circulation 38:15 1968
- 52 Myerberg R J Gelband H and Hoffman B F Functional characteristics of the gating mechanism in the canine A V conduction system Circ Res 28:136 1971
- 53 Damato A N Wit A L and Lau S H Observations on the mechanism of one type of so-called supernormal A V conduction Am Heart J 82:725 1971
- 54 Cranefield P F and Hoffman B F Conduction of the cardiac impulse II Summation and inhibition Circ Res 28:220 1971
- 55 Gertz G Kaplan H A Kaplan L G and Weinstein W Cardiac syncope due to paroxysms of ventricular flutter fibrillation and asystole in a patient with varying degrees of A V block and intraventricular block Am Heart J 16:225 1938
- 56 Watanabe Y A V conduction disturbances and electrophysiology Igaku No-Shoin 69:339 1967
- 57 Pick A and Katz L N Disturbances of impulse formation and conduction in the pre-excitation (WPW) syndrome—their bearing on its mechanism Am J Med 19:477 1955
- 58 James T N Morphology of the human atrioventricular node with remarks pertinent to its electrophysiology Am Heart J 62:156 1961
- 59 James T N The conducting pathways between the sinus node and A V node and between right and left atrium Am Heart J 66:498 1963
- 60 Lev M Anatomic considerations of anomalous A V pathways in Drefus L S and Likoff W editors Mechanisms and therapy of cardiac arrhythmias New York 1966 Grune & Stratton Inc p 665
- 61 Triex R C Anatomical considerations of the human atrioventricular junction in Drefus L S and Likoff W editors Mechanisms and therapy of cardiac arrhythmias New York 1966 Grune & Stratton Inc p 333
- 62 Durrer D Schuilenburg R M and Wellens H J J Preexcitation revisited Am J Cardiol 25:690 1970
- 63 Castillo C A and Castellanos A Jr The mechanism of ventricular preexcitation as studied by His bundle electrogram in Symposium on cardiac arrhythmias Denmark 1970 AB Astra p 329
- 64 Castellanos A Jr Castillo C A Acha A S and Tessier M His bundle electrograms in patients with short P R intervals narrow QRS complexes and paroxysmal tachycardias Circulation 43:667 1971
- 65 Coumel P Wajsbarger M Slama R and Bouvrain Y Intérêt de l'enregistrement de potentiels hisiens au cours du syndrome de Wolff Parkinson White à propos de six observations Acta Cardiol 26:188 1971
- 66 Lev M Lefler W B Langendorf B and Pick A Anatomic findings in a case of ventricular preexcitation (WPW) terminating in complete atrioventricular block Circulation 31:718 1966
- 67 Pick A and Fisch C Ventricular pre-excitation (WPW) in the presence of bundle branch block Am Heart J 55:504 1958
- 68 Pick A Aberrant ventricular conduction of escaped beats Preferential and accessory pathways in the A V junction Circulation 43:402 1956
- 69 Kistner A Atrioventricular junctional premature and escape beats with altered QRS and fusion Circulation 31:740 1966
- 70 Singer D H and Ten Eick R E Aberrancy Electrophysiologic aspects Am J Cardiol 28:381 1971
- 71 Rosenbaum M B Classification of ventricular

- extrasystoles according to form J Electrocardiol 2 289 1969
- 72 Wellens H Unusual occurrence of nonaberrant conduction in patients with atrial fibrillation and aberrant conduction AM HEART J 77:158 1969
- 73 Friedberg H D Mechanism of the Wedensky phenomenon in the left bundle branch Am J Cardiol 27 698 1971
- 74 Scherf D and Scharf M M Supernormal phase of intraventricular conduction AM HEART J 36 621 1948
- 75 Pick A and Fishman A P Observations in heart block Supernormality of AV and intra-ventricular conduction and ventricular parasystole under the influence of epinephrine Acta Cardiol 5 210 1950
- 76 Singer D H Lazzara R and Hoffman B P Interrelationships between automaticity and conduction in Purkinje fibers Circ. Res 21 537 1967
- 77 Rosenbaum M M Elzari M Lazzara J O Nau J N Levi R J and Halpern M S Intraventricular trifascicular blocks Review of the literature and classification AM HEART J 78 450 1969
- 8 Pick A Electrocardiographic features of exit block in Dreifus L S and Lakoff W editors Mechanisms and therapy of cardiac arrhythmias New York 1966 Grune & Stratton Inc p 469
- 9 Alan J Beritez D and Pilar G A functional discontinuity between the Purkinje and ventricular muscle cells Acta Physiol Latin Am 11 171 1961
- 80 Greenspan K Anderson G J and Fisch C Electrophysiologic correlate of exit block Am J Cardiol 28 197 1971
- 81 Myerberg R J Stewart J W and Hoffman H F Electrophysiologic properties of the canine peripheral AV conduction system Circ. Res 26 361 1970
- 8* Cranefield P F Klein H O and Hoffman B F Conduction of the cardiac impulse I Delay block and one way block in depressed Purkinje fibers Circ Res 28:199 1971
- 83 Wit A L Cranefield P F and Hoffman B F Slow conduction and reentry in the ventricular conduction system I Return extrasystole in canine Purkinje fibers II Single and sustained circus movement in networks of canine and bovine Purkinje fibers Circ. Res 30:11 11 1972
- 84 Goldenberg M and Rothberger C J Untersuchungen an der peripheren Muskulatur des Hundeherzens I Die Wirkungen unterschiedlicher Reize (angewendet auf die Parasystolie) Z Ges Exp Med 90 508 1933
- 85 Schaefer H Das Elektrokardiogramm Theorie und Klinik Heidelberg 1951 Springer Verlag
- 86 Scherf D and Schott A Extrasystoles and allied arrhythmias New York 1953 Grune & Stratton Inc.
- 87 Langendorf R and Pick A Parasystole with fixed coupling Circulation 33:304 1967
- 88 Roelandt J and Schamroth L Parasystolic ventricular tachycardia Observations on differential stimulus threshold as possible mechanism for exit block Br Heart J 34 505 1971
- 89 Watanabe Y Reassessment of parasystole AM HEART J 31 451 1971
- 90 Cohen H C Langendorf R and Pick A Mechanism of intermittent parasystole (Abstr.) Circulation 46 (Suppl II) 11 89 1972

Drug therapy of heart disease in pediatric patients

II The treatment of congestive heart failure in infants and children with digitalis preparations

Monika M Rutkowski, MD*

Sanford N Cohen, MD**

Eugenie F Doyle MD***

New York N Y

In general when drugs are administered to young pediatric patients the dosage regimen must be adjusted to take into account the immaturity of the various systems that control the amount of drug in the body at any time. Thus the usual rule is that infants require smaller doses than do older children and adults. However clinical experience has demonstrated that pediatric patients in congestive heart failure require higher doses of digitalis glycosides per unit weight than do adults.

The glycoside most commonly used to treat pediatric patients is digoxin. Its advantages are that it is readily absorbed, is available in convenient dosage forms, and is rapidly excreted from the body. In this paper we will summarize the data currently available concerning the factors which control the proper dosages of digoxin in infants and children and discuss the possible explanations for this seemingly paradoxical situation of higher digitalis requirement per unit weight in infants.

The pharmacokinetics of digitalis glyco-

sides could not be studied until recently, due to the difficulty of measuring them in biological fluids. Thus we are only now beginning to determine whether the apparent need for higher doses of digitalis in infants is due to poorer absorption, to more rapid excretion, or to greater end organ resistance.

Pharmacokinetics of digoxin

Absorption It is apparent from a study of the published data concerning the absorption of digoxin in infancy that there is little difference, if any, between the ability of pediatric patients to absorb this glycoside and for adults to absorb it. When serum levels of digoxin were studied by the radioimmunoassay technique it was apparent that infants receiving maintenance therapy orally were absorbing the drug as well as adults.^{1,2} Furthermore, in 20 infants aged 6 days through 5½ months, maximum blood levels of digoxin were achieved within 30 minutes after intramuscular administration and in 1 to 3 hours after oral adminis-

From the Departments of Pediatrics and Pharmacology, New York University School of Medicine. Received for publication March 22, 1973.

Reprint requests to: Dr. Monika M. Rutkowski, Instructor in Pediatrics, New York University School of Medicine, 550 1st Ave., New York, N.Y. 10016.

*Instructor in Pediatrics, New York University School of Medicine.

**Associate Professor of Pediatrics and Pharmacology, New York University School of Medicine. Physician in Charge of Nurseries, Bellevue Hospital. John and Mary R. Markle Scholar in Academic Medicine.

***Professor of Pediatrics and Director of Pediatric Cardiology, New York University Medical Center.

tration.⁶ This study was conducted by administering varying amounts of tritiated digoxin to patients in single doses.

Distribution The pharmacology of the digitalis glycosides has been studied extensively in adults through the use of tritiated digoxin.⁷ There was a constant relationship in mature individuals between tissue and serum digoxin concentrations with a ratio of 30 to 1 between the myocardium and the serum. In these studies the digoxin concentration was determined at a variable time after the administration of a single dose of digoxin. In a similar study 9 infants were given a single intravenous dose of digoxin and the tissue concentrations were measured after a variable amount of time had passed. The levels in these infants were claimed to be similar to those seen in adults,¹¹ although the actual value reported for adults is 52 ng per gram wet weight of heart,¹⁰ while the value reported for one 3 month old infant was 5.6 ng per gram wet weight of heart.⁸

In another study patients who were being maintained on digoxin therapy were evaluated with tritiated digoxin and similar values were reported for infants and adults. The concentration of digoxin per gram wet weight of heart reported by these investigators varied between 150 and 190 ng.⁴

The data which are available currently do not shed light upon the question of whether or not tissue digoxin concentration must be higher within the heart muscle for therapeutic effect in infants than in adults. However they imply that there can be a wide difference between the tissue concentration achieved after a single intravenous dose and that achieved after prolonged maintenance therapy. On the other hand it appears that the time required for distribution of a dose of digoxin is approximately the same in infants as in adults. Studies of the first phase of disappearance of digoxin from the circulation in patients of all ages indicate that the half time for distribution is between 0.5 and 1.5 hours.^{4, 8}

Excretion The mean half time for disappearance of a single dose of digoxin after this first phase (distribution) is approximately 34 hours in the adult (range 17 to 65 hours).⁸ In studies of a series of infants and young children the mean half time for disappearance of digoxin (after distribu-

tion) was found to be 32.5 hours (range 18 to 48 hours).⁴ These data are supported by the observation that 62.2 per cent of a dose of tritiated digoxin was recovered in the urine of adult patients within 3 days after administration⁷ and 55 per cent of a dose was recovered in a 3 day collection of urine from infants.⁸

In one study 5 infants with evidence of abnormal renal function (BUN elevations and abnormal inulin clearance) were included. The mean half time for elimination of digoxin in this group of patients was 50 hours (range 36 to 96 hours) and only 32 per cent of the dose was recovered in the urine after a 3 day collection.⁸ These data support the previous findings that patients with impaired renal function excrete digoxin more slowly than normals and should have their dose adjusted downward to prevent the build up of excessive serum and tissue concentrations.^{7, 11}

Mechanism of action

Most experimental studies on the mechanism of action of the cardiac glycosides have been conducted with ouabain. It is generally assumed that the results would be similar if digoxin were used.

There is no experimental evidence that suggests that the digitalis glycosides have a direct effect on myocardial contractile proteins. It is also apparent that the glycosides do not strengthen contractile force by influencing myocardial energy metabolism.

The glycosides probably act by facilitating the transmission of cell membrane excitation to the contractile elements within the cell.¹² It is thought that this effect is related to a specific inhibitory effect of the drug on the $\text{Na}^+ - \text{K}^+$ activated adenosine triphosphatase ($\text{Na}^+ - \text{K}^+$ ATPase) in the membrane. This inhibition probably leads indirectly to an increase in the free Ca^{++} concentration within the myocardial cell. This change in Ca^{++} concentration is thought to be the immediate trigger for the contractile process.¹³

Several investigators have studied the possibility that ouabain may effect myocardial $\text{Na}^+ - \text{K}^+$ ATPase differently at different ages. There was no age related difference in the $\text{Na}^+ - \text{K}^+$ ATPase activity of tissues from the hearts of 7 newborn lambs and 7 adult sheep. Furthermore there was

Table I Digoxin serum levels in infants and children on maintenance therapy

| Reference | Age | Mean digoxin dose (mg/kg) per 24 hrs | Mean digoxin serum level (ng/ml) |
|-----------|-----------|--------------------------------------|----------------------------------|
| 5 | 3-30 days | | |
| 2 3 | 1-11 mos | 0.018 | 1.8 |
| 5 | 1-12 mos | 0.021 | 2.7 |
| 2 3 | 2-14 yrs | 0.017 | 2.1 |
| 1 | 2-5 yrs | 0.015 | 1.4 |
| 1 | > 5 yrs | 0.016 | 1.4 |
| | | 0.009 | 0.9 |

Table II Relationship between digoxin dose, weight, surface area, and digoxin serum level in infants, children, and adults

| Reference | Age | Mean digoxin dose per 24 hrs | | A† | B† |
|-----------|-----------|------------------------------|---------------------|----------|----------|
| | | mg/kg* | mg/m ² * | | |
| 5 | 3-30 days | 0.018 | 0.35 | 1.7/1.5 | 3.3/1.5 |
| 2 3 | 1-11 mos | 0.021 | 0.36 | 1.8/2.0 | 3.9/2.0 |
| 5 | 1-12 mos | 0.017 | 0.33 | 1.6/1.6 | 3.1/1.6 |
| 2 3 | 2-14 yrs | 0.015 | 0.4 | 2.0/1.07 | 2.7/1.07 |
| 1 | 2-5 yrs | 0.016 | 0.4 | 2.0/1.07 | 3.0/1.07 |
| 1 | > 5 yrs | 0.009 | 0.27 | 1.3/0.7 | 1.6/0.7 |

*Values for average weight and surface area for the respective age group were assumed where not given in the references

† $\frac{\text{mg/M}^2 \text{ infant}}{\text{mg/M}^2 \text{ adult}}$ // $\frac{\text{ng/ml infant}}{\text{ng/ml adult}}$

† $\frac{\text{mg/kg infant}}{\text{mg/kg adult}}$ // $\frac{\text{ng/ml infant}}{\text{ng/ml adult}}$

no difference in ouabain binding activity, or in the stability of the binding complex between the tissues from lambs and those from sheep.¹³ On the other hand median myocardial Na-K ATPase activity was significantly greater in fetal and newborn puppies than in mature dogs.¹⁴ If these latter results are confirmed they may provide a partial explanation for the need of higher digoxin doses in infancy since they indicate that more drug may be needed to effect the same alteration in intracellular Ca⁺⁺ concentration in the immature myocardium.

Dosage recommendation

1 *Neonatal period* Newborn infants are more sensitive to digitalis than are older

infants. Thirty three per cent of 80 healthy newly born premature infants showed signs of digitalis intoxication when they were treated with digoxin 0.075 mg per kilogram of body weight.¹⁵ Only 9 per cent developed toxic signs when the dose was reduced to 0.050 mg per kilogram of body weight. Similarly bradycardia and atrioventricular block occur with greater frequency in newly born full term infants than in 2 to 4 month old infants given similar doses per unit of body weight.¹⁶ Therefore all neonates should be given lower doses per unit of body weight than 1 to 12 month old infants.

2 *Infancy* Several investigations have been conducted to examine the relationship of digitalizing and maintenance doses to

Table III Digoxin dosage in infants and children—mg/kg

| Route of administration | Newborn infant | Infants up to 1 yr | 1 to 2 years | Over 2 years | Maintenance dosage |
|-------------------------|----------------|--------------------|--------------|--------------|--------------------|
| Intramuscularly orally | 0.05 | 0.07 | 0.06 | 0.04 to 0.05 | 20 to 30% of TDD |
| Intravenously | 75% of above | 75% of above | 75% of above | 75% of above | |

Additional data: 1. 30 to 50 per cent of the TDD as a steady state plasma level 2. Digoxin is available in special form for pediatric use: pure (total digoxin) mg = 1 cc; (diluted form) 0.25 mg = 1 cc; (diluted) 0.05 mg = 1 cc.

serum concentration. They are summarized in Table I. Two studies were omitted from Table I: the first one because the results in the infant group differed considerably from those of other investigators and suggested that there is more rapid urinary excretion of the drug than has been noted by other workers¹⁷ and the second one because of the unusually high doses of digoxin used.⁴

It is apparent from the table that there were significantly higher serum levels in the infants than in the adults.^{2,3} The higher serum concentrations achieved in the infants correlate well with the higher doses administered to them if the dose is calculated per unit of body surface area rather than of body weight (Table II).⁴

In summary, infants absorb and excrete digoxin as well as adults. When higher doses are administered to infants they lead to correspondingly higher serum levels. There is no theory to explain the need for the higher dose in infancy satisfactorily. However, it is possible to speculate that if the heart of an infant has more myocardial Na-K ATPase activity than the heart of an adult, the infant might require more digoxin to inhibit this enzyme and produce a full therapeutic effect.¹⁸

3 Childhood. The serum concentration of digoxin was found to be the same in toddlers and older children as in adults.^{1,3} This is true despite the fact that the children received almost 100 per cent more digoxin per unit weight as maintenance as did the adults. The half time for elimination of digoxin from the body has only been determined in 3 children to date, but it appears to be similar to that found in infants.^{2,3} Further studies are therefore needed before we can understand the apparent discrep-

ancy between the dose of digoxin administered to children and the steady state serum concentration of the drug in them. It is possible that there is more rapid urinary excretion of digoxin in this age group or that the drug is bound and stored to a greater degree than in the infants. The tissue concentration of digoxin in a 7 year old child was measured and was found to be two to threefold greater than that found by the same investigators in a 3 month old infant.³ This observation points to the possibility that the latter theory may have some validity.

4 Dosage schedule. The dosage schedule presently recommended for digoxin therapy of pediatric patients of various ages is summarized in Table III. Note that the recommendation for a total digitalizing dose of digoxin in premature and full term newborn infants is 0.030 mg per kilogram of body weight. It rises to 0.070 mg per kilogram of body weight for infants up to 1 year of age and then decreases to 0.060 mg per kilogram of body weight for 1 to 2 year old infants. It decreases even further for children over the age of 2 years. The recommended dose for these patients is between 0.040 and 0.050 mg per kilogram of body weight. The maintenance regimen is calculated to replace 20 to 30 per cent of the digitalizing dose per day. If all of the digoxin is given intravenously the total dose should be only 75 per cent of the figures listed above.

Toxicity

Studies in experimental animals indicate that younger animals have a higher tolerance for digitalis glycosides than do adults of the same species. Thus the uniformly

lethal dose (LD_{100}) of glycoside for mature guinea pigs was only 75 per cent of the LD_{100} in newborns and the lethal dose for an isolated and continuously perfused adult Guinea pig heart was 50 per cent of the lethal dose for a heart from a 3 week old animal.¹⁸ Similarly the LD_{100} was significantly lower in adult dogs than in 15 to 65 day old puppies.¹⁹

It is not possible to determine the effect of age upon the toxicity of a dose of digoxin in the human from the published data. However, it is clear that even in man, toxicity is manifested at lower serum digoxin concentrations in adults than in infants. Signs of digitalis toxicity occur with great frequency in adults and children above the age of 2 years when the serum concentration exceeds 2 ng per milliliter. Levels below 2 ng per milliliter are well tolerated.^{7,17} It appears that in infants toxicity does not usually occur until the serum concentration exceeds 3.5 ng per milliliter, so long as acid base balance and electrolyte concentration are normal.^{2,25} It must be kept in mind, however that many of the early signs of digitalis intoxication (e.g., nausea, vomiting, visual disturbances) are difficult to perceive in infants.

The electrocardiographic signs of digitalis toxicity in infants are more likely to involve conduction impairment and supra-ventricular arrhythmia and not the ventricular arrhythmias seen in adults.

The following regimen is recommended for serious digoxin toxicity in infants and children:

- 1 Discontinue further digoxin
- 2 If the excessive dose was given intramuscularly, apply cold compresses and tourniquet to the extremity. If an excessive dose was administered by mouth gastric lavage should be attempted.
- 3 Potassium 0.05 mEq per kilogram of body weight should be given intravenously over 1 hour. The concentration of the solution in which it is administered should not exceed 80 mEq per liter. This dose may be repeated after 1 hour, if necessary.
- 4 Isoproterenol (1 vial = 200 mcg in 200 cc D₅W). Approximate dose up to 10 kilograms body weight 1 to 2 mcg per minute. 30 kilogram patient 2 mcg per minute. May be administered in cases of

complete A V block. External pacing may be necessary in such cases.

5 Diphenylhydantoin (2 to 3 mg per kilogram of body weight) may be administered intravenously in cases of tachyarrhythmias as a stat dose. A continuous infusion may be used if rapid rhythm recurs.

6 Lidocaine (1 mg per kilogram of body weight) propranolol (0.025 mg per kilogram of body weight) intravenously may be effective. This can be repeated X2 after five minute intervals, if no response occurs. Procainamide (3 to 5 mg per kilogram of body weight slowly intravenously) may be administered in cases of refractory tachycardias.

7 Cardioversion should be used if all other measures fail.

8 Blood should be drawn for determination of the concentration of digoxin.

Attempts to remove the drug by hemodialysis or by peritoneal dialysis have been unsuccessful and such procedures should not be relied upon in these cases.¹

REFERENCES

- 1 Krasula R W, Pellegrino P A, Hastreiter A R and Soyka L F. Serum levels of digoxin in infants and children. *Pediatrics* 81:566 1972.
- 2 Hayes C J, Gersony W M, Smith W B and Butler V P. Serum digoxin studies in infants and children. *Bull N Y Acad Med* 47:1227 1971.
- 3 Gersony W M, Hayes C J, Smith W B and Butler V P. Serum digoxin studies in infants and children. *Proceedings of the Society of Pediatric Research 41st Annual Meeting Atlantic City, New Jersey April 28 to May 1 1971*.
- 4 Murkin B L and Larese H. Immunoassay of serum digoxin levels in children: therapeutic implications. *Proceedings of the Society of Pediatric Research 41st Annual Meeting Atlantic City, New Jersey April 28 to May 1 1971*.
- 5 Rogers M C, Willerson J T, Coldblatt A and Smith T W. Serum digoxin concentrations in the human fetus, neonate and infant. *N Engl J Med* 287:1010 1972.
- 6 Hernandez A, Burton R M, Patakhan R D and Goldring D. Pharmacodynamics of H₂ Digoxin in infants. *Pediatrics* 44:418 1969.
- 7 Doherty J E. The clinical pharmacology of digitalis glycosides: a review. *Am J Med Sci* 253:382 1968.
- 8 Dungan W T, Doherty J E, Harvey C, Char F and Dalrymple G V. Tritiated digoxin. XVIII. Studies in infants and children. *Circulation* 46:983 1972.

- 9 Dunagan W T, Doherty J E and Harvey C. Studies with tritiated digoxin in infants (Abstract) *Circulation* 38 (Suppl VI) 68 1968
- 10 Doherty J E, Perkins W H and Flanagan W J. The distribution and concentration of tritiated digoxin in human tissues. *Ann Intern Med* 66:116 1967
- 11 Jelliffe R W and Blankenhorn D H. Improved method of digitalis therapy in patients with reduced renal function. *Circulation* 36 (Suppl II) 150 1967
- 12 Lee K S and Klaus W. The subcellular basis for the mechanism of inotropic action of cardiac glycosides. *Pharmacol Rev* 23 193 1971
- 13 Atwood G F and Dunkley S P. Maturity of the ouabain sensitive Na⁺/K⁺ ATPase receptor in the newborn lamb heart. *Circulation* 45 and 46 (Suppl II) 1136 1972
- 14 Miller W W and Cihland A. Developmental differences in canine myocardial adenosine triphosphatase activity. American Academy of Pediatrics 41st Annual Meeting Oct 14 and 15 1972 New York N Y
- 15 Levine O R and Blumenthal S. Digoxin dosage in premature infants. *Pediatrics* 29:18 1962
- 16 Marini A, Sereni F and Bottino D. Digoxin dosage in newborn animals and infants. *Pediatrics* 30 332 1962
- 17 Soyka L F. Clinical pharmacology of digoxin. *Ped Clin North Am* 19:241 1972
- 18 Wollenberger H, Jehl J A and Karsh M L. Influence of age on sensitivity of the Guinea pig and its myocardium to ouabain. *J Pharmacol Exp Ther* 108:152 1952
- 19 Halloran K M, Schimpff St C, Nicholas J G and Talner N S. Digitalis tolerance in young puppies. *Pediatrics* 46 730 1970

Annotations

Ischemic cardiomyopathy

The main function of a physician is to render high quality service to his patients. This he must do at a time when medical knowledge is extremely limited for most diseases anyway. For example among the frequently encountered cardiac illnesses are the cardiomyopathies the most resistant cardiac diseases to modern therapy. Therefore the most rewarding therapy for this group of illnesses is prevention. However regardless of all attempts to prevent cardiomyopathies patients develop these forms of cardiac disease and their physicians are challenged with management. After all the cardiomyopathies represent expressions of damage to the myocardium the source of pumping power for the circulation. And the blood must flow adequately or death ensues.

Accepting the term cardiomyopathy literally to mean heart muscle pathology the physician in America is confronted with myocardial ischemia due to coronary arterial obstructive disease as the most common cause of cardiac muscle pathology in America today. The late and readily recognized stage of the disease reveals all the symptoms and signs of the syndrome of cardiomyopathy such as congestive heart failure functional systolic murmurs (usually mitral and tricuspid) protodiastolic gallop rhythm and relatively high diastolic blood pressure in addition to the well known manifestations of coronary artery disease. The patients have evidence of myocardial ischemia such as that of angina pectoris without or with one or more myocardial infarcts. This classical syndrome of cardiomyopathy in patients with myocardial ischemia is often erroneously considered to be a cardiomyopathy due to other or unknown causes. However being aware that the syndrome can be produced by ischemic heart disease it is easy to recognize the cardiomyopathy as being due to myocardial ischemia and the patient can be properly and promptly treated accordingly. This syndrome I have chosen to call ischemic cardiomyopathy. Surely other cardiac diseases myocardial or otherwise may be superimposed upon it.

Patients with the late stages of ischemic cardiomyopathy require astute and meticulous management in order to achieve the best possible results. And many such patients must lead the life of a cardiac invalid or semi invalid if often necessary but unfortunately that this course is often necessary but the physician and the patient have no other choice and the sooner this way of life is thoroughly accepted by both the better the results and the happier the patient will be.

Finally it behooves the physician to recognize ischemic heart disease early as well as potential evidence of the disease before it develops in order to institute the best measures of prevention and management to interrupt or delay the onset of the late stages of ischemic cardiomyopathy. It should be realized that heart muscle pathology due to ischemia has a beginning and the beginning may be merely a slightly abnormal shape and/or magnitude of the T wave of the electrocardiogram in a patient who the physician decides from clinical data has potential or early coronary artery disease. Proper management of such patients immediately deliberately and adequately when first seen with angina pectoris or myocardial infarction is necessary for the best treatment and for prevention of advanced ischemic cardiomyopathy with its late and irreversible stages of the disease.

In the best interest of prevention of extensive and irreversible myocardial damage it must be remembered that cardiomyopathy can be extremely mild. The so-called classical syndrome reflects severe and irreversible damage to the myocardium. Therefore the diagnosis of ischemic cardiomyopathy should always be made when the diagnosis of ischemic heart disease is made. And when any disease exists in a patient which may result in coronary artery disease or when coronary artery disease is developing it behooves the physician to make the diagnosis of potential ischemic cardiomyopathy. The physician then should apply all measures available to him to reverse or arrest the coronary heart disease and/or at least reduce the work of the heart so that the myocardial demands for blood do not exceed the ability of the coronary circulation to satisfy those demands.

It is realized that some cardiologists limit the term cardiomyopathy to heart muscle disease of unknown cause. This practice is not to be ignored but it does restrict the physician and tend to confuse him from the more practical point of view of advising patients and it ultimately influences management of patients. A more detailed interpretation and classification of the cardiomyopathies is of greater value to the physician who is confronted daily with all sorts of diseases in his many patients.

G E Burch MD
Department of Medicine
School of Medicine
Tulane University
New Orleans 70112

The nitroblue-tetrazolium dye test and infection in the renal patient

The important observation that polymorphonuclear leukocytes are capable of spontaneously reducing the dye nitroblue tetrazolium (NBT) was made in 1968 by Park and co-workers.¹ The absolute number and percentage of circulating granulocytes reducing the dye increases during most bacterial infections and remains unchanged in the presence of viral infection. Since this initial report several large studies have confirmed the usefulness of this unstimulated NBT test in the detection and confirmation of bacterial infections in both children and adults.²⁻⁴ However there have been a paucity of studies on the usefulness of the NBT test in clinical conditions which carry a high risk of infection, one such area is renal disease. Specifically, there had been no studies on the efficacy of the NBT test in the differential diagnosis of infections in uremics or in immunosuppressed renal transplant patients. Consequently we undertook the following study.⁵

Two groups of renal patients were selected. The first group consisted of 32 uremic patients on whom NBT tests were performed randomly during their illness. Twenty seven of these patients were receiving chronic hemodialysis and none was receiving immunosuppressive therapy. The second group comprised ten post renal transplant patients 8 of whom were monitored serially with the NBT test following their transplants. We attempted to assess the usefulness of this test in the early diagnosis of bacterial infection and in its ability to differentiate infection from rejection. All of the transplant patients were receiving azathioprine (Imuran) at a uniform dosage of 2 mg per kilogram of body weight and steroids as Prednisolone at doses of 10 mg to 1 000 mg daily.

In our uremic group 23 of 32 patients were uninfected when NBT testing was performed. The mean value of NBT reduction in these patients was 5.1 per cent (i.e. 5.1 per cent of the polymorphonuclear leukocytes reduced NBT dye). The range was 0 to 15 per cent. The remaining nine patients in this group had documented infection 3 viral and 6 bacterial. The individuals with viral infection had normal NBT reduction (1 to 8 per cent) and of those with documented bacterial infection 5 of 6 had markedly elevated reduction of the dye (all greater than 40 per cent).

In our group of ten post transplant patients seven had proved bacterial infection at some time during their course. All of these showed significantly elevated NBT positivity (31 to 55 per cent) at times of infection. Four documented rejection episodes uncomplicated by concurrent infection occurred in 3 of the patients. During these episodes NBT reduction remained in the normal or unaffected range. One patient in the group of ten had an uncomplicated postoperative course and his NBT values remained normal on serial sampling. When Imuran and large doses of steroids (as much as one gram of Prednisolone daily) were administered we found

that the leukocytes of bacterially infected patients were capable of responding with increased dye reduction despite some previous suggestions to the contrary.⁶⁻⁸ The latter two studies measured NBT dye reduction in phagocytosing rather than in resting cells whereas our method quantified spontaneous dye reduction in non phagocytosing cells.

We were able to conclude from these findings that the NBT test is a valuable device for monitoring the infectious disease status of the renal patient. We base this conclusion on the following points: (1) Non infected uremic patient have normal baseline NBT values and in the face of bacterial infection are capable of mounting a positive NBT response. The test was also capable of distinguishing the viral from the bacterial infection despite similar clinical presentation. (2) Renal transplant patients are capable of responding to bacterial infections with a marked increase in the percentage of NBT positive cells. This is true in spite of the administration of large doses of steroids. (3) NBT test values remain normal in acute rejection allowing one to differentiate between bacterial infection and rejection even though the clinical presentations are so similar.

This study would also suggest by extrapolation that the NBT test is a valuable adjunct in following any patient whose underlying disease or therapy makes him especially susceptible to severe infection. Confirmation of the test's validity in any particular disease entity is a necessary prerequisite however.

Michael R. Wellman M.D.

Doris R. Miller M.D.

Department of Pediatrics

Division of Hematology

New York Hospital-Cornell Medical Center

New York, N.Y.

REFERENCES

1. Park B H, Fiksig S M, and Smithwick E M. Infection and nitroblue-tetrazolium reduction by neutrophils: A diagnostic aid. *Lancet* 2:532 1968.
2. Feigin R D, Shackelford P G, Choi S C, Flake K, E Franklyn F A, and Eisenberg C S. Nitroblue tetrazolium dye test as an aid in the differential diagnosis of febrile disorders. *J Pediatr* 81:230 1971.
3. Humbert J R, Marks M I, Hathaway W E, and Thoren C H. The histochemical nitroblue tetrazolium reduction test in the differential diagnosis of acute infections. *Pediatrics* 48:259 1971.
4. Matula G and Paterson P Y. Spontaneous in vitro reduction of nitroblue-tetrazolium by neutrophils of adult patients with bacterial infection. *N Engl J Med* 285:311 1971.
5. Wellman M R, David D S, Brennan B L.

- Lewy J E Stenzel K H Rubin A L and Miller D R The nitroblue tetrazolium test Usefulness in detecting bacterial infections in uraemic and immunosuppressed renal transplant patients *Lancet* 11:289 1972
- 6 Matula G and Paterson P Y NBT tests in a patient on steroids *Lancet* 1:803 1971
- 7 Miller D R and Kaplan H G Decreased

nitroblue tetrazolium dye reduction in the phagocytes of patients receiving Prednisone, *Pediatrics* 48:861 1970

- 8 Mandell G L Rubin W and Hook E W The effect of an NADH oxidase inhibitor (hydrocortisone) on polymorphonuclear leukocyte bactericidal activity *J Clin Invest* 49:1381 1970

The magnitude of risk of developing complete heart block in patients with LAD-RBBB

In recent years the electrocardiographic (ECG) as a marker of left axis deviation with right bundle branch block (LAD RBBB) has been recognized as a manifestation of bilateral bundle branch disease¹ and is a common forerunner of complete heart block.²⁻⁴ In spite of the great number of papers dealing with this topic that have appeared in the recent literature a question of the utmost importance still remains unanswered: what is the exact magnitude of the risk of development of complete heart block in patients with LAD RBBB? This risk has been estimated by previous workers (6 per cent⁵ 9 per cent¹⁰ 10 per cent⁶ and 13.6 per cent¹¹) but to our knowledge it has not been discussed so far in terms of duration of follow up.

The purpose of the present annotation is to report our experience in this matter. The ECG files of 50 patients who were seen initially at our hospital in 1968 or 1969 have recently been reviewed. Reasons for selecting these patients were as follows: All had an ECG which fulfilled previously described criteria for left anterior fascicular block combined with right bundle branch block.¹² None had evidence of atrioventricular conduction disturbance on the causal ECG taken at their first visit. Cases with myocardial infarction, congenital cardiac malformations and valvular heart disease were excluded. In our selected patients the underlying lesions responsible for the conduction disturbances were thought to be a sclerodegenerative process (Lenegre's or Lev's disease).¹³

The group under study consisted of 34 men and 16 women ranging in age from 43 to 83 years with an average age of 66 years. Several of these patients already had serial ECG tracings in their personal records when they were first examined in our hospital. This fact permitted a retrospective extension of the follow up periods. The patients were all repeatedly checked later and received particularly close scrutiny. At each examination they were carefully interviewed. If any symptom suggestive of paroxysmal atrioventricular block was discovered (episodes of dizziness, syncope or seizures) and if the ECG failed to disclose any atrioventricular conduction disturbance the patient was admitted to the intensive care unit and the ECG was continuously monitored for various periods of time up to 96

hours. Though tedious and expensive this procedure revealed rewarding findings in several instances.¹⁴

Altogether 16 patients (32 per cent) were discovered to present with transient or permanent complete A V block at some time in their illness. They were all treated by transvenous demand pacemakers. It should be stressed that in 8 of these 16 instances complete A V block was identified during the first 12 months of follow up. Actually these eight patients already had suggestive complaints at the time of their initial visit. Their presence in the series accounts for the skewing of the data towards abnormally high percentages of complete A V block of early onset. Therefore the most interesting data of this study are concerned with 38 patients of the series who had a follow up duration longer than 12 months and who had no dizziness or syncope when they were first seen. In the latter group follow up tracings were available for as long as 11 years with an average period of 4.8 years. To evaluate the long term prognosis of the association LAD RBBB our data were statistically treated using a method similar to the life table method proposed by Merrell and Shulman (1955).¹⁵ The expected incidence of A V block was calculated for each duration of follow up (Fig. 1). From the figures thus obtained it appears that the risk of developing complete heart block is about 6 per cent per year of follow up after recognition of LAD RBBB.

These results deserve some comments. To begin with the total incidence of complete A V block observed in our patients (21 per cent) appears somewhat higher than that quoted in previous reports.²⁻¹¹ Two reasons might account for this discrepancy. Firstly the mean duration of follow up in our series (4.8 years) was comparatively longer than in others (6.3 months¹⁰ and 19 months¹¹). Secondly the period under scrutiny started in 1968 at a time when one was already very eager to try and identify paroxysmal A V block in patients with LAD RBBB.

As predicted by Ranganathan and co-workers¹⁶ the risk of complete heart block appears extremely high when plotted with increasing time. This finding meets with the opinion expressed by Blondeau and Lenegre¹³ that if one excludes cases with myocardial

Probability of CHB (%)

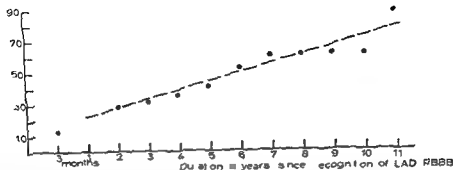


Fig 1 Risk of development of transient or permanent complete heart block in patients with LAD RBBB. The expected incidence of complete heart block was calculated for each duration of follow up using a method similar to that proposed by Merre and Shulman (1955).¹¹ If one only considers the portion of the diagram which covers (from 1 to 11 years of follow up (38 cases) the risk of developing complete heart block may be estimated at 6 per cent per year of follow up after identification of LAD RBBB ($y = 5.96x + 22.3$).

infarction and congenital cardiac malformations the natural outcome of patients with LAD RBBB is sooner or later towards complete A V block in approximately 100 per cent of the cases.

Nonetheless the duration in years that elapses between the first time LAD RBBB is recognized and the onset of complete A V block varies considerably in all published series. In some observations¹⁴ it was even as long as 23 years. This remark leads to a direct practical conclusion. At the present time any indiscriminate tendency to pace all patients with LAD RBBB as soon as this pattern is recognized appears unjustified. So far our policy in such cases has always been to await the first signs of paroxysmal A V block before implanting a cardiac pacemaker. This attitude admittedly debatable will possibly be reconsidered after completion of prospective studies of patients with LAD RBBB by means of his bundle electrography and atrial pacing. It is to be hoped that such investigations in progress in this institution as well as in many others will help identify at an early stage those patients who would be at greater risk for subsequent appearance of A V block and who might benefit from preventive pacing.^{14,15}

Henri E. Kulbertus M.D.
Chercheur Qualifié du FNRS

Division of Cardiology
Department of Medical Clinics and Semiology
University of Liège
School of Medicine
Liège Belgium

REFERENCES

1. Unger P N, Lesser M E, Hugel V H and Lev M. The concept of masquerading bundle branch block. An electrocardiographic-pathologic correlation. *Circulation* 17:397 1958.
2. Lénegre J. Contribution à l'étude des blocs de branche. Paris 1958. J. B. Baillière et fils. Editeurs.
3. Sugiura M, Ohada R, Hataoka, K. and Ohkawa S. Pathohistological studies on the conduction system in 14 cases of right bundle branch block associated with left axis deviation. *Jap Heart J* 10:121 1969.
4. Watt T B, Freud G E, Durrer D and Pruitt R D. Left anterior arborization block combined with right bundle branch block in canine and primate hearts. An electrocardiographic study. *Circ. Res* 22:57 1968.
5. Rosenbaum M, Elizari M V and Lazzari J O. Los Hemibloques. Buenos Aires 1968. Paidós Editores.
6. Lénegre J. Etiology and pathology of bilateral bundle branch block in relation to complete heart block. *Progr Cardiovasc Dis* 8:109 1964.
7. Lepeschkin E. The electrocardiographic diagnosis of bilateral bundle branch block in relation to complete heart block. *Progr Cardiovasc Dis* 6:445 1964.
8. Lasser R, Haft J and Friedberg Ch. Relationship of right bundle branch block and marked left axis deviation (with left parietal or paraventricular block) to complete heart block and syncope. *Circulation* 37:429 1968.
9. Kulbertus H and Collignon P. Association of right bundle branch block with left superior and inferior intraventricular block. Its relation to complete heart block and Adams-Stokes syndrome. *Br Heart J* 31:335 1969.
10. Watt T B Jr and Pruitt R D. Character cause and consequence of combined left axis deviation and right bundle branch block in human electrocardiograms. *Am Heart J* 77:460 1969.
11. Scanlon P J, Pryor E and Blount M G Jr. Right bundle branch block associated with left superior or inferior intraventricular block. Clinical setting, prognosis, relation to complete heart block. *Circulation* 42:1123 1970.
12. Kulbertus H, Collignon P and Humblet L. Left axis deviation and right bundle branch block. A vectorcardiographic study of 40 cases. 11th Symposium on Vectorcardiography. Proceedings of the Long Island Jewish Hospital.

- Lewy J E Stenzel K H Rubin A L and Miller D R The nitroblue tetrazolium test. Usefulness in detecting bacterial infections in uraemic and immunosuppressed renal transplant patients *Lancet* 11:289 1972
- 6 Matuli G and Paterson P Y NBT tests in a patient on steroids *Lancet* 1:803 1971
- 7 Miller D R and Kaplan H G Decreased nitroblue tetrazolium dye reduction in the phagocytes of patients receiving Prednisone *Pediatrics* 45:861 1970
- 8 Mandell G L Rubin W and Hook E W The effect of an NADH oxidase inhibitor (hydrocortisone) on polymorphonuclear leukocyte bactericidal activity *J Clin Invest* 49:1381 1970

The magnitude of risk of developing complete heart block in patients with LAD-RBBB

In recent years the electrocardiographic (ECG) association of left axis deviation with right bundle branch block (LAD RBBB) has been recognized as a manifestation of bilateral bundle branch disease^{1,2} and is a common forerunner of complete heart block.^{3,4} In spite of the great number of papers dealing with this topic that have appeared in the recent literature a question of the utmost importance still remains unanswered: what is the exact magnitude of the risk of development of complete heart block in patients with LAD RBBB? This risk has been estimated by previous workers (6 per cent⁵, 9 per cent⁶, 10 per cent⁷ and 13.6 per cent⁸) but to our knowledge it has not been discussed so far in terms of duration of follow up.

The purpose of the present annotation is to report our experience in this matter. The ECG files of 50 patients who were seen initially at our hospital in 1968 or 1969 have recently been reviewed. Reasons for selecting these patients were as follows: All had an ECG which fulfilled previously described criteria for left anterior fascicular block combined with right bundle branch block.⁹ None had evidence of atrioventricular conduction disturbance on the causal ECG taken at their first visit. Cases with myocardial infarction, congenital cardiac malformations and valvular heart disease were excluded. In our selected patients the underlying lesions responsible for the conduction disturbances were thought to be a sclerodegenerative process (Lenègre's or Lev's disease).¹⁰

The group under study consisted of 34 men and 16 women ranging in age from 43 to 83 years with an average age of 66 years. Several of these patients already had serial ECG tracings in their personal records when they were first examined in our hospital. This fact permitted a retrospective extension of the follow up periods. The patients were all repeatedly checked later and received particularly close scrutiny. At each examination they were carefully interviewed. If any symptom suggestive of paroxysmal atrioventricular block was discovered (episodes of dizziness, syncope or seizures) and if the ECG failed to disclose any atrioventricular conduction disturbance the patient was admitted to the intensive care unit and the ECG was continuously monitored for various periods of time up to 96

hours. Though tedious and expensive this procedure revealed rewarding findings in several instances.¹

Altogether 16 patients (32 per cent) were discovered to present with transient or permanent complete A V block at some time in their illness. They were all treated by transvenous demand pacemakers. It should be stressed that in 8 of these 16 instances complete A V block was identified during the first 12 months of follow up. Actually these eight patients already had suggestive complaints at the time of their initial visit. Their presence in the series accounts for the skewing of the data towards abnormally high percentages of complete A V block of early onset. Therefore the most interesting data of this study are concerned with 38 patients of the series who had a follow up duration longer than 12 months and who had no dizziness or syncope when they were first seen. In the latter group follow up tracings were available for as long as 11 years with an average period of 4.8 years. To evaluate the long term prognosis of the association LAD RBBB our data were statistically treated using a method similar to the life table method proposed by Merrel and Shulman (1955).¹¹ The expected incidence of A V block was calculated for each duration of follow up (Fig. 1). From the figures thus obtained it appears that the risk of developing complete heart block is about 6 per cent per year of follow up after recognition of LAD RBBB.

These results deserve some comments. To begin with the total incidence of complete A V block observed in our patients (21 per cent) appears somewhat higher than that quoted in previous reports.^{3,5,6,8} Two reasons might account for this discrepancy. Firstly the mean duration of follow up in our series (4.8 years) was comparatively longer than in others (6.3 months⁵ and 19 months⁸). Secondly the period under scrutiny started in 1968 at a time when one was already very eager to try and identify paroxysmal A V block in patients with LAD RBBB.

As predicted by Ranganathan and co-workers¹² the risk of complete heart block appears extremely high when plotted with increasing time. This finding meets with the opinion expressed by Blondecu and Lenègre¹³ that if one excludes cases with myocardial

Each subject was given 1 Gm sulfadimidine intravenously and at the same time an intravenous infusion of Inulin containing 10 to 25 Gm was started and continued for 4 hours. Blood and urine samples were taken before and 2, 3 and 4 hours after the injection of sulfadimidine and blood samples were taken 24 hours after the injection. Inulin was measured by Boyesen's method and sulfadimidine and N-acetylated sulfadimidine were estimated by the method of Bratton and Marshall.⁶

We found Inulin clearance as well as the mean clearances of sulfadimidine much lower in all uremic patients than in the controls.

Since only the non-protein bound sulfadimidine can be filtered through the glomeruli it was estimated by ultrafiltration through a cellophane membrane. Table I shows that although the total plasma concentrations of sulfadimidine are comparable in the two groups the non-protein bound sulfadimidine concentrations are higher in the uremic group. This accords with other findings.^{7,8} The explanation is not known. The lesser protein binding of sulfadimidine in uremia also favors the excretion of the drug in uremic patients; nevertheless the excretion of the drug is lower in uremic subjects than in controls.

Acetylated sulfadimidine constituted 43 per cent of the total plasma sulfadimidine in the uremic group and 21 per cent in the controls; for urine the figures were 38 per cent in the uremic patients and 52 per cent in the controls. Similar observations have been described.⁹ The higher percentage of acetylated sulfadimidine in plasma than in urine of uremic subjects indicates a lower renal excretion of acetylated sulfadimidine rather than an increased acetylation.

It has been shown that sulfadimidine in the human kidney is subjected to filtration and active tubular excretion as well as tubular reabsorption.⁵ Our data do not allow an estimation of any active secretion but the ratios between non-protein bound sulfadimidine clearance and Inulin clearance show that tubular reabsorption occurs in uremic as well as normal subjects. The parallel correspondence between Inulin clearance and non-protein bound sulfadimidine clearance however indicates that reduced filtration is mainly responsible for the decrease in sulfadimidine excretion in kidney failure. This explains too the finding that 24 hours after the injection of sulfadimidine between 7 and 23 mcg sulfadimidine per milliliter was still found in the

plasma of uremic subjects whereas none could be found in plasma from controls. This is in agreement with the demonstration of the 2 to 3 times longer plasma half-lives of sulfadimidine, methoxypyridazine sulfadimethoxine and Sulfadoxine in uremic subjects than in normal persons.¹ Our results also agree with those of Sharpstone⁴ who demonstrated reduced excretion of sulfamethoxazole as well as trimethoprim in kidney failure. On the other hand our findings are at variance with those of Williams and associates³ who explained for this discrepancy can be offered.

We conclude that there is a definite risk for an accumulation of the drug when uremic patients are treated with sulfadimidine.

Erna Fischer M.D.

Department of Medicine B and Biochemical Laboratory
Frederiksberg Central County Hospital
Hillerød, Denmark

REFERENCES

1. Kunitz C M. A guide to use of antibiotics in patients with renal disease. *Ann Intern Med* 67:151 1967.
2. Madsen S T and Larsen P F. Metabolic problems during treatment with long acting sulphonamides. *Proceeding of the Third International Congress of Chemotherapy* 1:644 1964.
3. Williams D M, Wimpey J and Asscher A W. Renal clearance of sodium sulphadimidine in normal and uremic subjects. *Lancet* 2:1058 1968.
4. Sharpstone P. The renal handling of trimethoprim and sulphamethoxazole in man. *Postgrad Med J* 45 (Suppl.):38 1969.
5. Boyesen E. A method for determination of inulin in plasma and urine. *Acta Med Scand* 266 (Suppl.):275 1952.
6. Bratton A C and Marshall E K. A new coupling component for sulfanilamide determination. *J Biol Chem* 128:537 1939.
7. Lunde P. Abnormal pharmacokinetics of phenytoin in a patient with uraemia. *Lancet* 2:832 1970.
8. Andreassen F. Protein binding of drugs in serum from patients with acute anuria. *Acta Pharmacol Toxicol* 29 (Suppl. 4):2 1971.
9. Weiner I M and Mudge G H. Renal tubular mechanisms for excretion of organic acids and bases. *Am J Med* 36:743 1964.

- Amsterdam 1971 North Holland Publishing Co p 272
- 13 Merrel M and Shulman L E Determination of prognosis in chronic disease illustrated by systemic lupus erythematosus *J Chronic Dis* 11:2 1955
- 14 Rangarathnam N Dhurandhar R Phillips J H and Wigle E D His bundle electrogram in bundle branch block, *Circulation* 45:282 1972

- 15 Blondeau M and Lenegre J Bloc atypique de la branche droite Paris 1970 Masson & Cie
- 16 Spurrell R A J Smithen C S and Sowton, E Study of right bundle branch block in association with either left anterior hemiblock or left posterior hemiblock using His bundle electrograms *Br Heart J* 34 800 1972

Renal excretion of sulfadimidine in normal and uremic subjects

The renal excretion rate of sulfonamides is an important factor in determining the concentrations of sulfonamides attainable in the body and in the urine. The restrictions imposed on antibacterial drugs in the treatment of patients with chronic renal failure have been described.¹ However, the renal excretion of sulfonamides in chronic renal failure has only been studied by a few workers.²⁻⁴

A higher renal clearance of sulfadimidine in uremic patients than in controls have been reported¹—a surprising result which could have important therapeutic implications. Since sulfonamides are still used in clinical practice and may even gain importance with the introduction of trimethoprim, we decided to compare the renal clearances of sulfadimidine and Inulin in 10 normal and 10 uremic per on

Table I Data on 10 uremic subjects and 10 controls

| Patient | Age (yr) | Sex | Body weight (kg) | Urine flow rate (ml/min) | Inulin clearance (ml/min) | Plasma sulfadimidine (μ g/ml) | | Total sul fadimidine urine con centration (μ g/ml) | Sulfadimidine clearance (ml/min) | |
|----------|-------------|-----|------------------------|--------------------------------|---------------------------------|--|---------|---|--|---------|
| | | | | | | Total | Unbound | | Bound | Unbound |
| Uremic* | | | | | | | | | | |
| 1 | 69 | F | 70 | 1.7 | 5 | 27 | 10.5 | 31 | 2.0 | 5.0 |
| 2 | 68 | M | 90 | 2.0 | 5 | 19 | 7.4 | 11 | 1.2 | 3.0 |
| 3 | 61 | M | 74 | 1.9 | 6 | 16 | 6.2 | 12 | 1.4 | 3.7 |
| 4 | 56 | F | 62 | 2.4 | 7 | 48 | 18.7 | 32 | 1.6 | 4.1 |
| 5 | 70 | F | 55 | 0.7 | 9 | 56 | 21.8 | 85 | 1.1 | 2.8 |
| 6 | 67 | M | 48 | 0.7 | 10 | 31 | 12.1 | 145 | 3.2 | 8.4 |
| 7 | 74 | F | 50 | 0.9 | 14 | 42 | 16.4 | 80 | 1.7 | 4.4 |
| 8 | 46 | F | 46 | 4.3 | 23 | 35 | 13.7 | 22 | 3.0 | 6.9 |
| 9 | 53 | M | 62 | 4.7 | 33 | 37 | 14.4 | 28 | 3.6 | 9.1 |
| 10 | 26 | M | 79 | 4.5 | 49 | 45 | 17.6 | 52 | 5.2 | 13.3 |
| Controls | | | | | | | | | | |
| 1 | 46 | M | 71 | 10.0 | 131 | 66 | 9.1 | 36 | 5.5 | 36.4 |
| 2 | 27 | F | 57 | 3.6 | 100 | 36 | 5.4 | 73 | 7.3 | 48.7 |
| 3 | 33 | M | 84 | 6.5 | 122 | 47 | 7.1 | 61 | 8.4 | 56.0 |
| 4 | 43 | F | 65 | 5.9 | 123 | 30 | 4.5 | 61 | 12.0 | 19.9 |
| 5 | 40 | M | 65 | 3.2 | 115 | 35 | 5.3 | 91 | 8.3 | 51.9 |
| 6 | 32 | F | 54 | 5.9 | 110 | 46 | 6.9 | 75 | 9.6 | 64.1 |
| 7 | 23 | M | 71 | 8.3 | 122 | 50 | 7.5 | 49 | 8.1 | 54.2 |
| 8 | 27 | F | 48 | 8.7 | 112 | 27 | 4.1 | 32 | 10.3 | 67.9 |
| 9 | 27 | F | 72 | 5.5 | 125 | 46 | 6.9 | 86 | 10.3 | 68.5 |
| 10 | 26 | M | 74 | 7.2 | 122 | 54 | 8.1 | 41 | 5.5 | 36.4 |

*Patients 1 and 3 and 8 to 10 had chronic glomerulonephritis and patients 2 and 4 and 7 had chronic pyelonephritis.

retically it is believed that all visceral functions can be brought under voluntary control by prolonged yogic training but perhaps their most fascinating claim has been the ability to stop the heart at will. However in most instances where this has been investigated so far it has turned out to be an exaggerated Valsalva maneuver in some form which makes the pulse and heart sounds imperceptible while the heart continues to beat at a slow rate.^{1,2}

Recently we had the rare opportunity of investigating an altogether different and very interesting demonstration of this supposed yogic control over the heart. Yogi Satyamurti, a sparsely built man of about 60 years of age, remained confined in a small underground pit for 8 days in what according to him was a state of Samadhi or deep meditation with all bodily activity cut down to the barest minimum. The pit was a 1.5 meter cube dug out in an open lawn surrounded by the Medical Institute buildings and was completely sealed from the top by bricks and cement mortar. The Yogi squatted on the floor of the pit with nothing on except a light cotton garment. About 5 liters of water was placed in one corner presumably for drinking but according to the Yogi only for keeping the air fully humid. An ECG (Lead II) was continuously monitored during these 8 days and various other laboratory investigations were carried out before and after. The ECG leads were kept short enough not to allow any free movement inside the pit.

The 12 lead ECG recorded before closing the pit was within normal limits (Fig 1 strip A) but a significant sinus tachycardia developed soon after. It increased progressively reaching a heart rate of 250 per minute on the second day (Fig 1 strip B). At 5.15 P.M. on the second day when the Yogi had been in the pit for about 29 hours to our great surprise a straight line replaced the ECG tracing (Fig 1 strip C). There was no electrical disturbance of any sort even at higher amplification and with different leads. There had been no slowing of the heart or signs of ischemia preceding this.

The straight line on the ECG persisted till the eighth morning. Then to our astonishment electrical activity returned about half an hour before the pit was scheduled to be opened. After some initial disturbance a normal configuration appeared. Although some sinus tachycardia was still there there was no other significant abnormality (Fig 1 strip D). The Yogi had informed us beforehand that he would begin to come out of his deep trance or suspended animation after nearly 7 days much in the same way that a normal person wakes up after a few hours sleep.

When the pit was opened on the eighth day the Yogi was found sitting in the same posture. One of us immediately went in to examine him. He was in a stuporous condition and was very cold (oral temperature was 34.8°C). On being taken out of the pit he developed severe shivering and this persisted for nearly 2 hours. A 12 lead ECG repeated in the laboratory subsequently was again within normal limits (Fig 1 strip E).

The Yogi and his admirers felt more than satisfied at this unequivocally documented proof of a remarkable yogic feat while we were left rather perplexed and confused. We were expecting some bradycardia

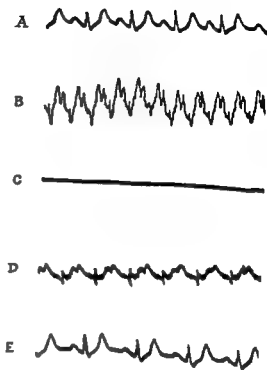


Fig 1 ECG tracing of the Yogi Lead II (Panel A) first day at 11.00 A.M. one hour before closing the pit. Normal sinus rhythm. Heart rate was 106 per minute. PR interval was 0.16 second. QRS interval was 0.08 second and there was an upright T wave. (Panel B) second day at 5.00 P.M. Severe sinus tachycardia with a heart rate of 250 per minute. a PR interval of 0.12 second. a QRS interval of 0.06 second and a high uptake of the ST segment. (Panel C) second day at 6.00 P.M. There is a straight line with no electrical disturbance. This continued for the next 5 days. (Panel D) eighth day at 7.30 A.M. half an hour before opening the pit. Sinus tachycardia is in evidence with a heart rate of 142 per minute. slightly prolonged PR interval of 0.20 second. a QRS interval of 0.06 second and an upright T wave. (Panel E) eighth day at 10.00 A.M. two hours after coming out of the pit. The tracing is normal with a heart rate of 98 per minute. PR interval of 0.16 second. a QRS interval of 0.08 second and an upright T wave.

and possible signs of myocardial ischemia but contrary to this there was severe tachycardia followed by a complete disappearance of all complexes. Any instrumental failure was ruled out by thoroughly checking the machine and also by the spontaneous reappearance of the ECG on the last day. A disconnection of the leads by the Yogi, quite a likely explanation ought to have given rise to considerable electrical disturbance but there was hardly any. Later on we tried all sorts of manipulations with the leads to simulate what the Yogi could have done

Letters to the Editor

Cardioversion after valve replacement

To the Editor

I read with interest the annotation by Drs Jenzer and Lown dealing with cardioversion after valve replacement (*Am Heart J* 81:810 1972). In reading it it became obvious that the authors have omitted reference to two critical issues. Unless such information is provided their comments and recommendations lose most of their relevance. These issues are:

1 How many of the 102 patients with valve replacement and the 58 patients with mitral valvotomy developed atrial fibrillation after the operation?

2 How many patients who were successfully cardioverted remained in sinus rhythm for more than two months?

We have demonstrated^{1,2} and others have amply confirmed our observations that postoperative atrial fibrillation and atrial fibrillation present prior to mitral valve surgery are two entirely different conditions. The former is an acute reaction to surgical trauma that bears no relation to the underlying disease (it occurs not only after cardiac but also after pulmonary operations) and is virtually 100 per cent convertible to sinus rhythm. The latter is a specific complication of mitral valve disease one that eventually leads to anatomic damage of atrial musculature.³ Our series showed a success rate in postoperative conversion of patients in the second category of 77 per cent but more than half of these lapsed back into atrial fibrillation after varying periods of time.

If an article describes the value of cardioversion and we recognize that cardioversion is good for oranges but only fair for apples should we know how many apples and oranges were mixed in the basket?

Arthur Selzer MD
Cardiology Division
Presbyterian Hospital
San Francisco Calif 94120

REFERENCES

- 1 Selzer et al. Treatment of atrial fibrillation after surgical repair of the mitral valve. *Annals Intern Med* 62:1213 1965
- 2 Popper R W, Knott J M S, Selzer A and Gerbode F. Arrhythmias after cardiac surgery I. Uncomplicated atrial septal defect. *Am Heart J* 61:455 1962
- 3 Bailey et al. Relation of left atrial pathology to atrial fibrillation in mitral valvular disease. *Annals Intern Med* 69:113 1968

Reply

To the Editor

Dr Selzer raises questions not germane to the intent of our brief communication. The objective was simply to provide evidence that valve replacement does not diminish the efficacy of cardioversion in restoring atrial fibrillation to sinus rhythm; furthermore it was to demonstrate that in these patients the procedure is not fraught with an increased incidence of immediate or late complications. If the physician therefore believes that sinus rhythm needs to be reestablished, cardioversion affords a simple, safe and effective method.

Of course Dr Selzer's questions can be factually answered. Of the 102 patients with valve replacement 19 developed atrial fibrillation for the first time after the operation. The success rate of cardioversion and the required energy were the same irrespective of when the arrhythmia first occurred. Our experience to date indicates that if atrial fibrillation has been present for less than one year prior to valvular surgery, the likelihood for sustained sinus rhythm for the ensuing year approximates 50 per cent.

We are of course well aware that sinus rhythm has a greater chance of being maintained in patients in whom atrial fibrillation follows rather than precedes valvular operation. The crucial question therefore is which patient following valve correction will maintain sinus rhythm after cardioversion when the arrhythmia has been established for some months. We are not aware of any clinical shibboleth to permit such distinction. In the absence of guide lines we believe that all such patients should be subjected to the relatively innocuous procedure of cardioversion. Whether the arrhythmia merely precedes or follows valvular operation is far too simple and inadequate a discriminant. Dr Selzer hails from a state well known for the abundance and variety of its fruit; he should therefore have no difficulty in distinguishing apples from oranges.

Hans Jenzer MD
Bernard Lown MD
Cardiovascular Laboratories
Harvard School of Public Health
Cardiology Division
Peter Bent Brigham Hospital
Boston, Mass 02115

The Yogic claim of voluntary control over the heart beat: an unusual demonstration*

To the Editor

Yogis in India have long been reputed to develop a remarkable control over bodily functions. Theo-

*A detailed report on the Yogis is appearing in *The Indian Journal of Medical Research* November 1973.

Table I Effectiveness of practolol in 145 patients

| Type of arrhythmia | No digitalis | | | Previous digitalis | | |
|---------------------|--------------|-------------|-----------|--------------------|-------------|-----------|
| | No | Good result | No result | No | Good result | No result |
| Sinus tachycardia | 10 | 9 | 1 | 21 | 18 | 3 |
| Atrial fibrillation | 21 | 15 | 6 | 25 | 24 | 1 |
| Atrial flutter | 22 | 10 | 12 | 14 | 12 | 2 |
| Atrial tachycardia | 17 | 12 | 5 | 15 | 14 | 1 |

2 Atrial fibrillation (7 patients) normal sinus rhythm (N.S.R.) in 1 slowing of the H.R. in 5 (average reduction = 35 per cent) no result in 1

3 Atrial flutter (4 patients) N.S.R. in 1 (after a short sequence of atrial fibrillation) slowing in 2 (average reduction = 32.5 per cent) no result in 1

4 Supraventricular tachycardia (3 patients) N.S.R. in 3 slowing in 1 (reduction = 38 per cent) no result in 1

5 Ventricular ectopics (5 patients) total disappearance in 1 decrease in 1 no result in 1 Good results (N.S.R. or H.R. below 100 beats per minute) were thus obtained in 88.4 per cent of the patients with supraventricular tachyarrhythmias. In none of these acutely ill patients did worsening of the respiratory status occur during or within 15 minutes after drug administration.

The antiarrhythmic effectiveness of practolol is markedly potentiated by previous digitalization (Table I). In 70 patients with supraventricular tachyarrhythmias of various etiology practolol used alone was effective in 46 (67 per cent). 24 patients (33 per cent) did not respond at all. In a similar group of 75 patients who had been previously digitalized good results were obtained in 68 (90.7 per cent). 7 patients only (9.3 per cent) did not respond at all.

We do feel as did Dr Fromgren that practolol may be used safely in patients with bronchospasm and that it is a drug of choice for the treatment of supraventricular tachyarrhythmias complicating their respiratory insufficiency.

Jean Pierre Van Durme M.D.

Leo Bossaert M.D.

Paul Vermeire M.D.

René Pannier M.D.

Department of Cardiology

Akademisch Ziekenhuis

University of Gent

9000 Gent-Belgium

REFERENCES

- 1 Van Durme J P Bossaert L Vermeire P and Pannier R. One hundred fifty cases of cardiac arrhythmias treated with practolol (ICI 50172). *Acta Cardiol Suppl* xv '85 1972
- 2 Van Durme J P Bossaert L Vermeire P and Pannier R. Le traitement des troubles du

rythme par le Practolol (Eraldin) G Ital Cardiol 21 161 1972

Right atrial electrocardiogram resulting in hemopericardium

To the Editor

In a recent communication Drs Homesley and Zelenik re-emphasized the potential hazards of routine investigations (*Am Heart J* 84 133 1972). We would like to stress again this point by reporting briefly a case of lethal hemopericardium which occurred as a result of a right atrial electrocardiogram (RAE). A 61 year old man was admitted to our department because of the recent onset of heart failure and supraventricular tachycardia. The RAE was recorded through a Vigon catheter to further investigate the arrhythmia. The procedure went uneventfully but seven hours later seizures occurred suddenly while blood pressure dropped. The patient became cold and cyanotic. Distention of the neck veins appeared. He was given 40 mg of metaraminol which failed to bring up the blood pressure. Three more seizures occurred within an hour. The patient became unconscious as a slow junctional rhythm was recorded. Attempts at resuscitation were ineffective. Pathological studies revealed a traumatic perforation of the right atrium wall which resulted in a fatal acute tamponade.

Robert Hoyat M.D.

Claude Seban M.D.

Richard Benaim M.D.

Paul Chiche M.D. F.A.C.C.

Service de Cardiologie et Urgences Circulatoires

Hopital Tenon

4 rue de la Chue

75020-Paris France

Atrial ectopic tachycardia

To the Editor

I read with much interest the article by Goldreyer Gallagher and Damato in a recent issue of the *Journal* (*Am Heart J* 83 205 1973). Again these excellent ongoing studies from their laboratory have demonstrated the utility of intracardiac recording techniques for the evaluation of potential electrophysiologic mechanisms of supraventricular

inside the pit (notwithstanding the total darkness and his ignorance of ECG technique) but in every case there was marked disturbance. Therefore although it is obviously difficult to believe that the Yogi could have completely stopped his heart or decreased its electrical activity below a recordable level, we still had no satisfactory explanation for the ECG tracings before us.

Apart from this, the Yogi had of course endured total starvation, sensory deprivation as well as the discomfort of a very humid, closed atmosphere for 8 days. We did not pay much attention to anoxia, thinking that sufficient ventilation could occur through the bare earth on the sides of the pit. The loss of weight (4.5 kilograms) and other biochemical changes were essentially the same as can be expected in starvation under similar conditions. They certainly discount any remarkable depression of the metabolic rate.

The more optimistic amongst us considered this feat to be a marvelous extension of the hypometabolic wakened state of yogic meditation, as described by Wallace and co-workers³ and the conditioned learning of autonomic responses in rats reported by DiCara.⁴ The skeptics, however, were inclined to take the whole thing as some cleverly disguised trick. But for the present, we only want to put this interesting experiment on record just as an intriguing and inconclusive attempt of a Yogi to demonstrate a voluntary control over his heart beat.

L. K. Kothari M.Sc. M.A.M.S.

Reader in Physiology

Arun Bordia M.D.

Reader in Cardiology

O. P. Gupta M.D.

Professor of Pathology

Ravindra Nath Tagore Medical College
and Hospital Udaipur, India

Reprint requests to L. K. Kothari M.Sc. M.A.M.S.

REFERENCES

1. Anand B. K. and Chhina G. S. Investigations on yogis claiming to stop their heart beats. *Indian J. Med. Res.* 19:90 1961.
2. Wenger M. A., Bagchi H. K. and Anand B. K. Experiments in India on voluntary control of the heart and pulse. *Circulation* 24:1319 1961.
3. Wallace R. K., Benson H. and Wilson A. F. Wakened hypometabolic physiologic state. *Am. J. Physiol.* 221:795 1971.
4. DiCara L. V. Learning in the autonomic nervous system. *Sci. Am.* 222:31 1970.

Atropine in acute myocardial infarction

To the Editor

The discussion of the value of atropine in acute myocardial infarction at the last meeting of the American College of Cardiology in San Francisco (Medical World News, March 9, 1973) is the reason for the following comments.

The risk of giving atropine in patients with coronary disease has been demonstrated many years ago by Wayne and Laplace.¹ They found in patients with angina pectoris that after administration of atropine a lesser physical exertion elicited anginal pain. Scherf and Schnabel² demonstrated that following an intravenous injection of 2 mg of atropine in patients with syphilitic aortitis and angina pectoris due to stenosis of the orifice of a coronary artery, anginal pain with marked depression of the RS-T segments appeared. The increased rate induced by atropine in many (not all) patients with coronary disease is responsible for these results. This is illustrated by another observation reported by the same authors: A diabetic patient died several minutes after the intravenous injection of 1/4 mg of atropine. The autopsy revealed severe arteriosclerosis of both coronary arteries.

Because the appearance of a marked increase of rate following an injection of atropine cannot be predicted, I consider this medication risky in patients with coronary disease.

David Scherf, M.D.

Dept. of Medicine

New York Medical College

1249 Fifth Ave.

New York, N.Y. 10019

REFERENCES

1. Wayne E. J. and Laplace L. B. Observations on angina of effort. *Clin. Sci.* 11:103 1933.
2. Scherf D. and Schnabel P. Atropin bei Angina pectoris. *Klin. Wochenschr.* 13:1397 1934.

Practolol in treating tachyarrhythmias

To the Editor

We read with great interest the article of Dr H. Formgren entitled "Practolol in the treatment of tachyarrhythmias in patients with bronchial asthma" (*Am. Heart J.* 84:710 1972). We would however like to take issue with his statement that "Practolol has not been previously given to asthmatics for treatment of concomitant cardiovascular disorder."

Since 1969 we have indeed been using practolol for the treatment of acute tachyarrhythmias and have reported our results previously.¹ We would therefore like to report briefly our experience in patients with bronchospasm due to asthma, bronchiale or chronic obstructive lung disease.

When assessing the antiarrhythmic effectiveness of a drug we feel that it always should be given intravenously and under continuous electrocardiographic recording. Practolol was given as follows: 5 mg every minute until a therapeutic result was obtained or a total dose of 25 mg was reached; if necessary a maintenance dose of 100 to 200 mg twice daily was given.

The results obtained in 29 patients hospitalized with acute respiratory failure can be summarized as follows:

1. Lowering of the heart rate (HR) below 100 beats per minute (average reduction = 31.7 per cent) in 10 patients with sinus tachy.

Book reviews

ECHOCARDIOGRAPHY By Harvey Feigenbaum M D and Sonia Cheng B.S Philadelphia Pa 1972 Lea & Febiger 239 pp

Feigenbaum has rendered an extremely valuable service to cardiology in this lucid and practical publication on echocardiography The book is well written and well organized The presentations are designed for beginners and the subjects discussed are extremely well selected This book presents very well the present state of clinical echocardiography a rapidly changing field in cardiology All cardiologist internists and students should study this book and learn echocardiography now and continue their education from the medical literature This excellent well written book by an authority is highly recommended

TREATMENT OF HEART DISEASE IN THE ADULT 2nd ed By Ira Lloyd Rubin M D Harry Gross M D and Sidney M Arbeit M D (Clinical Pharmacology) Duncan E Hutcheon M D Philadelphia Pa 1972 Lea & Febiger 506 pp Price \$27 00

This second edition on the treatment of heart disease in the adult is a good one The presentations are clear and the subjects of practical nature It contains 70 chapters concerned with coronary heart disease drugs hypotensive heart disease and the other common forms of heart disease

The authors also discuss endocarditis pericarditis arrhythmias pacemakers digitalis pregnancy and heart disease and other clinical and therapeutic problems related to heart disease The presentations are good The authors lucidly define their therapeutic approaches to heart disease This is a practical and valuable book for practicing physicians as well as for physicians in training

ON PULSATILE AND STEADY ARTERIAL FLOW THE GTS CONTRIBUTION By A Iberall M Cardon and E Young Upper Darby Pa 1973 General Technical Services Inc 225 pp Price \$6 00

This small book on pulsatile and steady arterial blood flow emanates from the research of the NASA program on space medicine The contributors have considered the important factors influencing arterial blood flow from mathematical and engineering points of view These six essays should interest physiologists rheologists and investigators of the peripheral circulation The authors of the essays have attempted to consider quantitatively the variables that influence arterial flow Their considerations are thought provoking but still fail to define entirely the extremely dynamic changes in the many complex variables that modify the circulation of blood in arteries This is a very good and important publication

arrhythmias. This article emphasizes that in addition to re entry ectopic atrial pacemakers may be responsible for paroxysmal supraventricular arrhythmias. Furthermore they stress that atrial arrhythmias due to re entry generally require (1) atrial premature depolarizations and (2) AV nodal conduction delay. Recent reports however have commented on the fact that paroxysmal re entrant atrial arrhythmias may also occur because of (1) progressive AV nodal conduction delay without atrial premature beats and (2) progressive delay in conduction in the distal conduction system.^{2,3} These reports would I am sure add further support to comments that all re entrant atrial tachycardias may not have the same mechanism.

W J Mandel MD
Cedars Sinus Medical Center Los Angeles Calif
J Lozano MD
Hospital Central de las Fuerzas Armadas
Caracas Venezuela
H Hayakawa MD
Asippon Medical School Tokyo Japan

REFERENCES

- 1 Goldreyer B N, Gallagher J J and Damato A N The electrophysiologic demonstration of atrial ectopic tachycardia in man. *AM HEART J* 85:205 1973
- 2 Mandel W J, Kermuer A I, Blum R L and Hayakawa H Critical AV conduction prolongation as the initiating mechanism in re entrant tachycardia: a study using His bundle recordings. *J Electrocardiology* 3:39 1972
- 3 Lozano J, Mandel W J, Hayakawa H, Shine K I and Eber L M Reentrant tachycardia: participation of the distal A V conduction system. *Chest* 63:23 1972

Reply

To the Editor

Drs Mandel, Lozano and Hayakawa correctly point out that not all atrial tachycardias are due to re entry and in fact that was the purpose of our recent report.¹ Those atrial tachycardias which are due to re entry may be due to re entry in the AV node,¹ the distal AV conduction system and possibly the sinoatrial node. Our experience however in studying more than 30 patients with classical paroxysmal supraventricular tachycardia has shown the mechanism of the tachyarrhythmia to be due to AV nodal re entry in each case. Although other sites of re entry are possible, I would stress that AV nodal re entry is the typical mechanism of paroxysmal supraventricular tachycardia.

Bruce N Goldreyer MD
Recipient Samuel Beilet Professorship in Cardiology
Hospital of the University of Pennsylvania
Cardiac Catheterization Laboratory
3400 Spruce Street
Philadelphia Pa 19104

REFERENCES

- 1 Goldreyer B N, Gallagher J J and Damato A N The electrophysiologic demonstration of

atrial ectopic tachycardia in man. *AM HEART J* 85:205 1973

- 2 Goldreyer B N and Damato A N The essential role of atrial ventricular conduction delay in the initiation of paroxysmal supraventricular tachycardia. *Circulation* 48:619 1971

Systolic time intervals in man

To the Editor

Contrary to the statement of Dr Lindquist and associates,¹ we did not correct the PEP for heart rate in our paper.²

Jean M Pouget MD
Willard S Harris MD
Department of Medicine
University of Illinois College
of Medicine and West Side
Veterans Administration Hospital
Chicago Ill

REFERENCES

- 1 Lindquist V A, Spangler R D and Blount S G Jr A comparison between the effects of dynamic and isometric exercise as evaluated by the systolic time intervals in normal man. *AM HEART J* 85:227 1973
- 2 Pouget J M, Harris W S, Mayron H R and Naughton J P Abnormal responses of the systolic time intervals to exercise in patients with angina pectoris. *Circulation* 43:289 1971

Reply

To the Editor

We apologize for this error in our paper¹ and note that your data² recorded only the QS₂ and LVET corrected for heart rate. Had the PEP been similarly corrected by using the regression equations of Weissler and colleagues,³ your normal subjects would have demonstrated a mean decrease of 14 msec with exercise. Hence our observations on the significance of protocol and method of heart rate correction for the STI in general are still valid.

Valdemar Lindquist MD MRCP
Richard D Spangler MD FCCP
S Gilbert Blount Jr MD
Department of Medicine
Division of Cardiology
University of Colorado Medical Center
Denver Colo 80720

REFERENCES

- 1 Lindquist V A, Spangler R D and Blount S G Jr A comparison between the effects of dynamic and isometric exercise as evaluated by the systolic time intervals in normal man. *AM HEART J* 85:227 1973
- 2 Pouget J M, Harris W S, Mayron H R and Naughton J P Abnormal responses of the systolic time intervals to exercise in patients with angina pectoris. *Circulation* 43:289 1971
- 3 Weissler A M, Harris W S and Schoenfeld C D Systolic time intervals in heart failure in man. *Circulation* 37:149 1968

Book reviews

ECHOCARDIOGRAPHY By Harvey Feigenbaum M D and Sonin Cheng BS Philadelphia Pa 1972 Lea & Febiger 239 pp

Feigenbaum has rendered an extremely valuable service to cardiology in this lucid and practical publication on echocardiography. The book is well written and well organized. The presentations are designed for beginners and the subjects discussed are extremely well selected. This book presents very well the present state of clinical echocardiography, a rapidly changing field in cardiology. All cardiologists, internists and students should study this book and learn echocardiography now and continue their education from the medical literature. This excellent well written book by an authority is highly recommended.

TREATMENT OF HEART DISEASE IN THE ADULT 2nd ed. By Ira Lloyd Rubin M D, Harry Gross M D and Sidney R. Arbeit M D (Clinical Pharmacology). Duncan E. Hutcheon M D Philadelphia Pa 1972 Lea & Febiger 506 pp Price \$27.00

This second edition on the treatment of heart disease in the adult is a good one. The presentations are clear and the subjects of practical nature. It contains 20 chapters concerned with coronary heart disease, drug, hypotensive heart disease and the other common forms of heart disease.

The authors also discuss endocarditis, pericarditis, arrhythmias, pacemakers, digitalis, pregnancy and heart disease and other clinical and therapeutic problems related to heart disease. The presentations are good. The author lucidly defines their therapeutic approaches to heart disease. This is a practical and valuable book for practicing physicians as well as for physicians in training.

ON PULSATILE AND STEADY ARTERIAL FLOW: THE CTS CONTRIBUTION By A. Iberall, M. Cardon and E. Young Upper Darby Pa 1973 General Technical Services Inc. 225 pp Price \$6.00

This small book on pulsatile and steady arterial blood flow emanates from the research of the NASA program on space medicine. The contributors have considered the important factors influencing arterial blood flow from mathematical and engineering points of view. These six essays should interest physiologists, rheologists and investigators of the peripheral circulation. The authors of the essays have attempted to consider quantitatively the variables that influence arterial flow. Their considerations are thought provoking but still fail to define entirely the extremely dynamic changes in the many complex variables that modify the circulation of blood in arteries. This is a very good and important publication.

Books received

ASTHMA: A PRACTICAL GUIDE FOR PHYSICIANS Prepared by a joint committee of the Allergy Foundation of America and the American Thoracic Society. Published by the National Tuberculosis and Respiratory Disease Association in cooperation with the Allergy Foundation of America. 1973. 72 pages.

LEARNING HOW TO LIVE WITH HEART TROUBLE By Arthur J. Snider. Chicago, Illinois. 1973. Budlong Press Company. 125 pages. Price \$1.75.

DEVELOPMENTS IN BIOMEDICAL ENGINEERING Edited by Martin M. Black. New York. 1973. Crane, Russak & Company, Inc. 972 pages. Price \$19.75.

CARING FOR THE DYING PATIENT AND HIS FAMILY: A MODEL FOR MEDICAL EDUCATION—MEDICAL CENTER CONFERENCES Edited by Dr. Austin H. Kutscher and Michael R. Goldberg. New York. 1973. Health Sciences Publishing Corporation. 72 pages.

PHYSIOLOGY AND BIOPHYSICS OF DIGESTION, METABOLISM, ENDOCRINE FUNCTION AND REPRODUCTION 20th ed. Edited by Theodore C. Ruth, Ph.D. and Harry D. Patton, Ph.D. M.D. Philadelphia. 1973. W.B. Saunders Company. 391 pages.

THE ALLIED HEALTH PROFESSIONAL AND THE PATIENT: TECHNIQUES OF EFFECTIVE INTERACTION By Ruth Purtilo, R.P.T. Philadelphia. 1973. W.B. Saunders Company. 229 pages.

THE CRISIS TEAM—A HANDBOOK FOR THE MENTAL HEALTH PROFESSIONAL By Julian Lieb, M.B.B.Ch., Ian I. Lipsitch, M.D., and Andrew Edmund Slaby, M.D. Hagerstown, Maryland. 1973. Harper & Row Publishers, Inc. 186 pages. Price \$6.95.

✓ **PROGRESS IN CARDIOLOGY, No. 2** Edited by Paul V. M.D. and John F. Goodwin, M.D. Philadelphia. 1973. Lea & Febiger Publishers. 290 pages. Price \$15.00.

✓ **ESSENTIALS OF PEDIATRIC CARDIOLOGY** By James H. Moller, M.D. Philadelphia. 1973. F.A. Davis Company. 124 pages. Price \$4.50.

✓ **THE MANAGEMENT OF NEONATES AND INFANTS WITH CONGENITAL HEART DISEASE** By Donald M. Billig, M.D. and Marshall B. Kreidberg, M.D. New York. 1973. Grune & Stratton, Inc. 182 pages. Price \$12.75.

✓ **ADVANCES IN EXPERIMENTAL MEDICINE AND BIOLOGY, vol. 26: PHARMACOLOGICAL CONTROL OF LIPID METABOLISM** Edited by William L. Holmes, Rodolfo Paoletti, and David Kritchevsky. Proceeding of the Fourth International Symposium on Drugs Affecting Lipid Metabolism held in Philadelphia, Pennsylvania, September 8-11, 1971. New York, London. 1972. Plenum Press. 359 pages. Price \$22.50.

THE HARVARD FATIGUE LABORATORY: ITS HISTORY AND CONTRIBUTIONS By Steven M. Horvath and Elizabeth C. Horvath. Englewood Cliffs, New Jersey. 1973. Prentice Hall, Inc. 182 pages. Price \$9.95.

✓ **THE SILENT DISEASE: HYPERTENSION** By Lawrence Galton. New York. 1973. Crown Publishers, Inc. 210 pages. Price \$5.95.

Editorial

His bundle electrocardiography

Charles Fisch MD
Douglas P Zipes MD
Indianapolis Ind

All new techniques introduced into medicine are ultimately judged by their contribution to our understanding of pathophysiology; enhancement of diagnostic ability; insight into the mechanism of therapeutic interventions; and to the overall management of the ill patient.

The technique of His bundle electrocardiography (HBE) although relatively new has interested many investigators and has resulted in a number of published studies with considerable unanimity of findings. For this reason one can attempt to evaluate even at this early stage of the development the contribution of HBE to our understanding of the mechanism of arrhythmias and to the management of the patient with heart disease. This editorial was written with a clear understanding that information inherent in HBE is far from complete and that much of it is yet to be realized.

The technique of HBE involves passing a catheter across the tricuspid valve which requires sophisticated and expensive instrumentation and considerable technical skill. However in experienced hands the technical aspects of the procedure are

relatively simple and safe. An electrode catheter is introduced percutaneously into the femoral vein with its tip positioned near the septal leaflet of the tricuspid valve. The His potential (H) appears as a well defined most often bipolar spike between the low atrial (A) and ventricular (V) electrograms. The interval between the earliest onset of the surface P wave or a high right atrial deflection and the low right atrial deflection (P-A) is a measure of intra-atrial conduction. The interval between A and H (A-H) is a measurement of the conduction across the A-V node and varies in duration from 50 to 120 msec depending on the cycle length and autonomic influences. The interval from H to V (H-V) is a measure of His Purkinje conduction time and is determined by the interval between the His deflection and the earliest ventricular activity recorded in any lead. The H-V interval is a measure of conduction through the His bundle distal to the recording electrode; the bundle branches and the Purkinje system up to the point of ventricular activation. In contrast to a relatively wide range of values for the A-H the H-V interval is fairly constant measuring 30 to 50

From the Division of Cardiology, Indiana University School of Medicine, Indianapolis, Ind. Received for publication July 11, 1973. Accepted for publication August 1, 1973. Reprint requests: Dr. Charles Fisch, Division of Cardiology, Indiana University School of Medicine, 635 North Walnut Street, Indianapolis, Ind. 46202.

msec with an average value of 45 msec.

The ability to separate A-V nodal and His-Purkinje (H-P) conduction has enhanced our understanding of normal and abnormal atrioventricular (A-V) conduction. Abnormal A-V conduction may be due to prolongation of P-A, A-H, H-V, or all three. In patients with a P-R interval in the upper limits of normal, the H-V may be abnormally prolonged and when subjected to atrial pacing may reveal latent block of the H-P system and thus aid in the differential diagnosis of syncope. It should be pointed out, however, that the H-V may be normal and the patient may still have periodic complete distal His block. HBL confirmed the fact that when the QRS duration is normal, the A-V block is usually in the A-V node. On the other hand, in the presence of a prolonged QRS, the block may be either in the A-H or in the H-V regions or in both. Second degree block of the Wenckebach type is most often, but not always, A-V nodal in origin, while Type II is nearly always due to a block in the H-V system. Concealed discharge of the bundle of His is a cause of unexpected first or second degree A-V block, had been proposed from analysis of the surface ECG and conclusively proved with HBL in the human. Incomplete penetration of the A-V node or bundle of His (concealed conduction) has been repeatedly demonstrated in the human with HBE.

HBL clearly confirmed pre-excitation of the ventricles in the Wolff-Parkinson-White (WPW) abnormality. In this syndrome, the A-H is normal and the H-V is shortened. With atrial pacing, the A-H lengthens normally, but the A-V conduction by way of the bypass remains unchanged and consequently the H-V shortens, the His deflection eventually becoming lost in or actually following ventricular excitation. The syndrome of short P-R, normal QRS and tachycardia has been shown in some instances to be due to foreshortening of the A-H interval without any change of the H-V period. The exact reason for shortening of the A-H conduction remains obscure. In some of the reported cases, the response of the A-H interval to pacing is qualitatively the same as in the normal but quantitatively much less pronounced. Our understanding of the possible interrelationship of the electro-

physiologic manifestations of ventricular pre-excitation with anatomic correlates (James-Mahaim or Kent bypass) must await further studies. The mechanism of integrade block with preservation of retrograde conduction has been the subject of considerable discussion. HBE clearly demonstrates an H potential between the ventricular and atrial electrogram, thus strongly supporting unidirectional block with preservation of retrograde A-V conduction.

Although the major contribution of the His recording has been to our understanding of A-V conduction and its various aberrations, HBE has provided some help with analysis of ectopic rhythms. For example, HBE demonstrated clearly that many supraventricular tachycardias are due to re-entry within the A-V node and that accelerated idioventricular rhythm, so frequently seen in acute myocardial infarction, is ventricular in origin. The ever present problem of differential diagnosis of supraventricular aberrancy and ventricular ectopy can be resolved with HBE recording. In the former, H precedes V with a normal H-V time, while in the latter the H deflection is often lost within the ventricular electrogram or if it precedes the local ventricular electrogram, it follows the earliest moment of ventricular activation recorded in the multiple surface leads. Rarely, ventricular ectopic impulse will activate the bundle of His before giving rise to a QRS and in such a situation an H-V shorter than normal will be recorded. There is always the possibility, however remote, that one simply fails to record the H spike during supraventricular rhythm and in erroneous diagnosis of ventricular tachycardia may be entertained.

There is little doubt from the foregoing brief review that the HBE has and will continue to contribute to our understanding of arrhythmias and that this avenue of investigation should be pursued by those studying mechanisms of arrhythmias and who are thus equipped for routine HBE. On the other hand, the utility of HBE in everyday routine care of patients with heart disease is much less secure and in fact is yet to be defined.

Irrespective of the invasive nature of the procedure, the technique and documentation require

the HBE in the management of the patient will hinge on two factors namely (1) evidence that HBE offers considerable advantage over a carefully analyzed surface ECC correlated with the total clinical picture and equally important (2) our ultimate ability to utilize fully the information obtained with the HBE in the actual care of the patient

At present we are unable to benefit fully from the information inherent in the HBL because complementary parallel clinical facts are not available. The following two examples will serve to illustrate this point.

The recognition from the HBE that a given supraventricular tachycardia is reentrant in origin rather than automatic will have little effect on the therapy of the arrhythmia until such time that drugs specific for one or the other of the two mechanisms becomes available. Similarly, recognition that a prolonged H V time is the cause of the prolonged P R interval can be helpful only if the long term prognosis of the H V delay is known.

Because of the instrumentation and technique involved in recording His bundle activity the surface ECC will remain the method of choice or perhaps the only method for diagnosis of cardiac arrhythmias available to the vast majority of physicians. In fact in emergent situations the lack of HBL should not even raise doubts in the physician's mind as to whether or not proper treatment will be administered. The availability of cardioversion a nonspecific therapeutic intervention makes it possible after a careful consideration of the total clinical picture to treat the occasional patient with an undiagnosed life threatening arrhythmia without the necessity of having to identify the exact mechanism or site of origin of the arrhythmia. However such a nonspecific therapeutic approach to arrhythmias may not always be possible nor desirable. For example patients with arrhythmias suspected to be due to a ventricular aneurysm and considered for resection must have the arrhythmia properly diagnosed. In such an instance the HBL may prove to be essential and the patient can be referred to a laboratory designed for such an investigation. In elective situations in which the patients can benefit from HBL referral for such studies may be desirable.

Records of the last 40 patients who have undergone HBE in our institution were reviewed for contribution of this technique to the management of these patients. The results from each study were analyzed to determine the potential benefit of HBE to the clinical care of given individuals. In only five instances did the HBL prove to be of significant help by providing information not available after analysis of the surface ECC. In four cases it aided in the differential diagnosis of ventricular and supraventricular arrhythmia with aberrancy and in one case it identified distal His block in a patient with syncope.

HBE has confirmed a number of concepts which evolved from elegant analysis of surface cardiograms. It has generated some new knowledge and will continue to enhance our understanding of arrhythmias. To date however its contribution to the management of patients with heart disease has been limited. This is due not only to (1) the fact that much of the clinically needed information can be obtained from the surface ECC but also in some measure to (2) deficiencies in our knowledge of the basic mechanisms specific therapy and prognosis of various arrhythmias. This information if and when available will make the data derived from HBL clinically more useful. However until such parallel information does become available the HBL will remain of limited value as a clinical tool.

REFERENCES

- 1 Scherlag H J, Lazzarini H, Helfant R H, Stein E, Berkowitz W D and Damato A N. Catheter technique for recording His bundle activity in man. *Circulation* 39:13, 1969.
- 2 Damato A N, Gallagher J J and Lazzarini H. Aplanctus of His bundle recordings in diagnosis of conduction disorders. *Tex Heart J* 18:601, 1972.
- 3 Narula O S, Scherlag B J, Simet I and Javier R E. Atrioventricular block localization and classification by His bundle recording. *Am J Med* 40:146, 1971.
- 4 Roers K M, Rabinowitz S H and Gunnar R M. Pseudo AV block secondary to premature non-pyramidal His bundle depolarization demonstrated by His bundle electrocardiography. *Circulation* 42:167, 1970.
- 5 Caldwell A Jr, Cuthbert C A, Agha A S, et al. Torsades de pointes in patients with heart block: interval narrow QRS complexes and paroxysmal tachycardia. *Circulation* 43:667, 1971.

Stress electrocardiography in the evaluation of aortocoronary bypass surgery

Arthur Dodge MD*

Donald G Kassebaum MD**

Herbert J Griswold MD***

Portland, Oreg

The aortocoronary vein bypass operation has pre-empted other myocardial revascularization techniques because of the directness of approach and the high rate of symptomatic improvement.¹⁻⁹ Moreover functional improvement has been documented in terms of significantly increased myocardial blood flow^{2,5,8,10,12} and improved ventricular performance.^{14,15}

Despite early symptomatic and functional improvement it is still not clear whether saphenous vein grafts will remain patent over longer periods of time and offset the relentless progression of coronary arteriosclerosis. Furthermore the psychological effect of surgery is large in patients with angina pectoris, and any revascularization operation has been shown to decrease symptoms in a significant number of cases. For these reasons objective and easily repeatable methods of evaluation are required to determine if myocardial is-

chemia is reduced after aortocoronary bypass.

Stress electrocardiography is a practical non-invasive, safe and reproducible procedure which has been shown previously to correlate closely with the angiographic results of myocardial revascularization.¹⁶ This report describes evaluation of saphenous vein aortocoronary bypass by standardized pre and postoperative hypoxemia and graded exercise electrocardiography. In this study the results of stress electrocardiography are correlated with the symptomatic response and angiographic evidence of graft patency.

Methods

Patients Details of the clinical material are given in Tables I to III. Forty six patients having saphenous vein aortocoronary bypass surgery between July 1970 and January 1972 were studied.

From the Division of Cardiology, Department of Medicine, University of Oregon Medical School, Portland, Oregon. Supported by Program Project Grant HL 06336 and Graduate Training Grant HL 05791 of the National Heart and Lung Institute.

Received for publication Oct. 17, 1972.

Reprint requests to Donald G. Kassebaum, MD, Division of Cardiology, University of Oregon Medical School, Portland, Oregon 97201.

† Doctoral Clinical Cardiology Trainee, Division of Cardiology, University of Oregon Medical School, Portland, Oregon.

* Professor of Medicine (Cardiology) and Chief of Medicine, Multnomah Hospital, University of Oregon Medical School, Portland, Oregon.

*** Professor of Medicine and Head, Division of Cardiology, University of Oregon Medical School, Portland, Oregon.

Table 1 Clinical data and stress ECG results

| | Those having postoperative angiography | | All patients |
|---|--|---------------|----------------------|
| | One or both grafts open | No open graft | |
| Number of patient | 34 | 7 | 45 |
| Male | 32 | 5 | 41 |
| Female | 2 | 2 | 4 |
| Age (yr) range (average) | 38-61 (50) | 42-55 (46) | 38-61 (50) |
| Angina pectoris | | | |
| Duration (yr) range (average) | 1-13 (5) | 0-5-10 (4) | 0-5-18 (5) |
| Functional Class | | | |
| 2 | 3 | 0 | 3 |
| 3 | 29 | 0 | 38 |
| 4 | 2 | 1 | 3 |
| Prior myocardial infarction | 74 | 5 | 79 |
| Time since infarction (yr) range (average) | 0-5-13 (4) | 1-8 (3) | 0-5-13 (4) |
| Preoperative stress ECG | | | |
| range (average) | | | |
| Final Sao_2 (%) | 66-89 (77) | 67-90 (77) | 66-90 (77) |
| Maximum work (kpm/min) | 100-1 000 (388) | 700-750 (314) | 100-1 000 (380) |
| Maximum heart rate | 88-150 (109) | 94-141 (110) | 88-150 (109) |
| Postoperative stress ECG | | | |
| range (average) | | | |
| Final Sao_2 (%) | 67-90 (77) | 67-88 (79) | 67-90 (78) |
| Maximum work (kpm/min) | 100-800 (515) | 150-800 (379) | 100-800 (503) |
| Maximum heart rate | 100-150 (127) | 100-143 (118) | 100-165 (126) |
| Difference from preoperatively average (significance) | | | |
| Hypoxemia (% Sao_2) | 0 (NS) | +2 (NS) | +1 (NS) |
| Maximum work (kpm/min) | +127 ($p < 0.01$) | +65 (NS) | +123 ($p < 0.001$) |
| Maximum heart rate | +18 ($p < 0.001$) | +8 (NS) | +17 ($p < 0.001$) |
| Time of postoperative | | | |
| range (average) | | | |
| Stress ECG | 3-14 (6) | 3-11 (6) | 3-14 (6) |
| Angiography | 1-12 (5) | 3-6 (4) | 1-12 (5) |

prospectively. There was one operative death; this report describes the results in the 45 survivors. All 45 had pre and post operative stress electrocardiograms and preoperative coronary arteriography. 41 had postoperative coronary arteriograms. All patients were Functional Class 3 or 4 (New York Heart Association) except for three who were Functional Class 2. All had chronic disabling angina pectoris and none had hypertension, left ventricular hypertrophy, congestive heart failure, atrial fibrillation or were taking digitalis glycosides. Twenty-nine patients had prior myocardial infarction. Postoperative stress electrocardiography was done between 3 and 14 months averaging 6 months after surgery. In most cases it was performed

the day before postoperative arteriography so that both the patient and the investigators were unaware of the status of the graft. The ages ranged from 38 to 61 years averaging 51 years. There were 41 men and 4 women.

Surgery. Twenty-six patients had single saphenous vein bypass grafts. In 14 cases the graft was implanted to the right coronary artery (RCA); in 12 others it was implanted to the left anterior descending coronary artery (LAD). One patient had triple grafts. Fifteen patients had double grafts. Of the 19 with two or more grafts, 17 had grafts to both the RCA and LAD and two had grafts to both the LAD and circumflex artery.

Clinical evaluation. A standardized ques-

Table 11 Stress TCG and angiographic findings in patients with aortocoronary bypass grafts

| Patient | Age sex | Preoperative resting ECG | Preoperative coronary arteriography | | | Postoperative angiography | | | |
|------------------------|------------|--------------------------------|--|------|------|---------------------------|---------------------------------------|------------------------------------|-----|
| | | | | | | Time (mo) | Coronary changes from preoperative | Location and condition of graft | |
| | | | RC† | LAD | C | | | | |
| GROUP I | | | | | | | | | |
| 1 L M | 46 M | AMI | 40 | 100 | 0 | 4 | None | LAD | |
| 2 H B | 61 M | N | 100* | 100* | 0 | | | | |
| 3 L D | 50 M | Ischemia | 100* | 100 | 80 | 4 | None | LAD RC | C |
| 4 C W | 60 M | AMI | 100 | 100* | 90* | 1 | None | LAD | C |
| 5 A D | 60 M | N | 100* | 80 | 50 | 9 | Collat to RC | LAD | RC |
| 6 D I | 38 M | N | 90* | 50* | 0 | 12 | None | LAD RC | |
| 7 I M | 51 M | Ischemia | 0 | 95 | 100* | 5 | None | LAD C | |
| 8 S G | 48 M | IMI | 100 | 80* | 50 | 4 | None | LAD | |
| 9 T W | 61 M | N | 100* | 80* | 0 | | | | |
| 10 H I | 52 M | AMI | 50* | 100* | 40 | 6 | Ocel proximal RC | RC | LAD |
| 11 W M | 45 M | AMI | 75* | 95 | 50 | 9 | None | LAD RC | |
| 12 W M | 39 M | AMI IMI | 100* | 90* | 0 | 5 | None | LAD RC | |
| 13 T M | 54 M | IMI | 100* | 100* | 90 | 4 | None | LAD RC | |
| 14 T W | 51 M | N | 100* | 70* | 50 | 4 | None | LAD C | |
| Mean | | | | | | | | | |
| Standard error of mean | | | | | | | | | |
| Significance (p) | | | | | | | | | |
| GROUP II | | | | | | | | | |
| 1 I K | 51 M | IMI | 95* | 40 | 0 | 1 | None | RC | |
| 2 T W | 52 M | IMI | 100* | 100 | 0 | 5 | None | RC | |
| 3 D I | 43 M | N | 80* | 0 | 0 | 1 | None | RC | |
| 4 C M | 50 M | IVCD | 90* | 50* | 90* | 7 | Ocel C | LAD RC | |
| 5 M I | 59 M | N | 70* | 0 | 90 | 5 | Ocel C | RC | |
| 6 C B | 42 M | N | 0 | 100 | 0 | 5 | None | LAD | |
| 7 W C | 58 M | IVCD | 90* | 50 | 75 | 9 | Ocel C RC | RC | |
| 8 C W | 51 M | Ischemia | 100* | 70 | 50 | | | | |
| 9 L C | 41 M | IMI | 100* | 80* | 40 | 6 | None | LAD RC | |
| 10 C W | 61 M | N | 100* | 75* | 0 | 4 | 90% stenosis LAD | LAD | |
| Mean | | | | | | | | | |
| Standard error of mean | | | | | | | | | |
| Significance (p) | | | | | | | | | |

*Grafted artery

†Abbreviations: RC = right coronary; LAD = left anterior descending; C = circumflex; N = normal; AMI = anterior myocardial infarction; NS = not significant

tionnaire was completed each time the patient was seen quantitating the activity level, frequency of angina pectoris and the consumption of nitroglycerin tablets. The clinical result was classified as excellent when the patient was able to return to work and/or had no angina pectoris. Patients with a good clinical result were able to more than double their physical activity or had less than half the preoperative frequency of angina and use of nitroglycerin tablets, or both. A fair result was charac-

terized by postoperative increase in activity up to twice the preoperative level or a third as much angina or nitroglycerin usage.

Angiography. Selective coronary, left ventricular and aortic (graft) angiography was done by the percutaneous transfemoral technique of Judkins²⁰ using 35 mm cineangiography and sequential cut films. The extent of disease was expressed as the percent narrowing of the lumen of each vessel.

| Preoperative stress ECG test | | | | | Postoperative stress ECG test | | | | | Clinical improvement |
|------------------------------|-------------------------------------|------------|--------------------|----------------|-------------------------------|-------------------------------------|------------|--------------------|----------------|----------------------|
| Hypoxemia | | Exercise | | | Hypoxemia | | Exercise | | | |
| Pos or neg | Max hypoxemia (% SaO ₂) | Pos or neg | Max work (kpm/min) | Max heart rate | Pos or neg | Max hypoxemia (% SaO ₂) | Pos or neg | Max work (kpm/min) | Max heart rate | |
| + | 0 | — | 400 | 108 | — | 77 | — | 800 | 150 | Excellent |
| + | 83 | + | 400 | 100 | — | 83 | — | 600 | 110 | Excellent |
| — | 83 | + | 400 | 115 | — | 80 | — | 600 | 120 | Excellent |
| + | 85 | — | 600 | 97 | — | 85 | — | 800 | 150 | Excellent |
| + | 80 | + | 300 | 94 | — | 10 | — | 300 | 135 | Excellent |
| + | 75 | — | 350 | 136 | — | 78 | — | 350 | 136 | Good |
| + | 66 | + | 400 | 115 | — | 67 | — | 600 | 142 | Good |
| + | 80 | + | 400 | 115 | — | 83 | — | 600 | 136 | Fair |
| + | 78 | + | 400 | 120 | — | 84 | — | 800 | 165 | Excellent |
| + | 18 | — | 400 | 120 | — | 74 | — | 600 | 137 | Excellent |
| + | 78 | + | 200 | 100 | — | 73 | — | 600 | 115 | Excellent |
| + | 78 | + | 600 | 111 | — | 72 | — | 600 | 116 | Excellent |
| + | 89 | + | 200 | 130 | — | 75 | — | 600 | 140 | Excellent |
| + | 81 | + | 400 | 95 | — | 80 | — | 400 | 105 | Good |
| | 79 | | 389 | 111 | | 77 | | 589 | 134 | |
| | 16 | | 32 | 36 | | 15 | | 43 | 46 | |
| | | | | | | NS | | <0.001 | <0.001 | |
| — | 74 | — | 200 | 88 | — | 73 | — | 600 | 125 | Good |
| — | 84 | — | 400 | 107 | — | 79 | — | 600 | 120 | Excellent |
| — | 73 | — | 400 | 120 | — | 73 | — | 600 | 150 | Excellent |
| — | 80 | — | 400 | 111 | — | 81 | — | 200 | 100 | Good |
| — | 78 | — | 400 | 100 | — | 68 | — | 600 | 140 | Good |
| — | 68 | — | 400 | 95 | — | 10 | — | 800 | 136 | Excellent |
| — | 72 | — | 800 | 105 | — | 74 | — | 400 | 100 | Good |
| — | 16 | — | 500 | 100 | — | 81 | — | 700 | 120 | Good |
| — | 87 | — | 300 | 107 | — | 87 | — | 800 | 150 | Excellent |
| — | 84 | — | 200 | 84 | — | 82 | — | 400 | 107 | Excellent |
| | 76 | | 400 | 102 | | 76 | | 570 | 125 | |
| | 21 | | 57 | 36 | | 18 | | 63 | 63 | |
| | | | | | | NS | | <0.05 | <0.005 | |

IM1 = 1 so myocardial infarction ICD = 1 true 1 d et del y C Max = 11 c r l Oct = oc Max = maximum

Stress electrocardiography Patients were studied in the postabsorptive state and no one was taking digitalis at this time or for the two weeks preceding the test. In the hypoxemia test patients inhaled a gas mixture of 10 per cent oxygen and 90 per cent nitrogen and the level of hypoxemia was monitored constantly by an earpiece oximeter (Waters) calibrated against Van Slyke arterial oxygen determinations. The test was terminated if chest pain occurred if at least 0.5 mm segmental ST depression

appeared in the electrocardiogram (ECG) or if the arterial oxygen saturation fell below 70 per cent. The maximum duration of hypoxemia was 20 minutes.

Graded supine exercise was performed by pedaling a bicycle ergometer (Elema Shonander) in the method standardized in this laboratory.¹⁰ The first level of exercise was 100 to 200 kpm per minute; this was increased by increments of 100 to 200 kpm per minute until at least 1.0 mm segmental ST depression occurred in the

Table III Stress LCG and angiographic findings in patients with aortocoronary bypass grafts

| Patient | Age sex | Preoperative resting ECG | Preoperative coronary arteriography | | | Postoperative angiography | | | | |
|------------------------|------------|--------------------------------|--|------|-----|---------------------------|---------------------------------------|------------------------------------|---------|--|
| | | | | | | Time (mo) | Coronary changes from preoperative | Location and condition of graft | | |
| | | | RC† | LAD | C | | | Open | Closed | |
| GROUP III | | | | | | | | | | |
| 1 I H | 50 I | N | 90* | 90* | 0 | 1 | 95% occl LAD less collat | I AD RC | | |
| 2 H B | 42 M | N | 0 | 100* | 0 | 6 | None | | I AD | |
| 3 H G | 42 M | N | 40 | 100* | 0 | 3 | None | | LAD | |
| 4 G M | 55 M | IMI | 100* | 50 | 0 | 2 | None | | RC | |
| 5 R C | 38 M | IMI AMI | 100 | 100 | 50 | 1 | None | LAD | | |
| 6 R N | 53 M | IMI | 90 | 50* | 100 | 5 | Occl OM | LAD | | |
| 7 R R | 47 I | IMI | 100 | 50 | 50 | 4 | None | | RC | |
| 8 J H | 58 M | IMI | 100 | 50 | 80 | 6 | 70% steno in I AD distal to graft | LAD | | |
| 9 H H | 39 M | N | 0 | 80* | 0 | 6 | None | LAD | | |
| 10 H S | 43 M | IMI | 100* | 100 | 100 | 3 | None | | I AD RC | |
| 11 R D | 49 M | N | 100 | 80* | 0 | 6 | None | | I AD | |
| 12 F H | 61 I | N | 100* | 100 | 60 | 2 | None | RC | | |
| 13 J C | 47 F | N | 100* | 80 | 90 | 5 | None | | IC | |
| 14 W L | 45 M | IVCD | 100* | 100* | 90 | 5 | 95% steno in RC | I AD RC | | |
| 15 W D | 49 M | N | 100* | 0 | 0 | 4 | None | RC | | |
| 16 M J | 56 M | N | 0 | 90* | 0 | 3 | None | I AD | | |
| 17 O P | 54 M | N | 100* | 50 | 0 | | | | | |
| 18 I D | 58 M | AMI | 100* | 90* | 100 | 6 | None | LAD | IC | |
| Mean | | | | | | | | | | |
| Standard error of mean | | | | | | | | | | |
| Significance (p) | | | | | | | | | | |
| GROUP IV | | | | | | | | | | |
| 1 K N | 56 M | N | 90* | 95 | 0 | 4 | Occl RC | RC | | |
| 2 V B | 59 M | N | 80* | 30 | 95 | 3 | Occl RC C collat to C | RC | | |
| 3 A M | 46 M | AMI | 65* | 100* | 90 | 4 | None | LAD RC | | |

*Grafted artery

Abbreviations: RC = right coronary; IAD = left anterior descending; C = circumflex; OM = obtuse marginal; N = normal; AMI = acute myocardial infarction; Max = maximum; NS = not significant

ECG or chest pain, dyspnea or fatigue terminated the exercise. All patients exercised at a given level for 3 minutes and rested for 3 minutes between exercise periods.

A standard 12 lead LCG was obtained from each patient prior to stress. Special skin electrodes were attached to the extremities and six chest positions, and the ECG was obtained by a Hewlett Packard (Sanborn Series 350) 4 channel photographic recorder at a paper speed of 50 mm per second. A switching network made it possible to register a complete 12 lead

ECG in 10 to 15 seconds. Any selected group of 4 leads was displayed continuously on a Hewlett Packard 4 channel oscilloscope (Sanborn Model 569A) and a cardiographometer (Parks Electronics Laboratory) coupled to the electrocardiograph provided continuous indication of the heart rate. The gain of the electrocardiograph was adjusted for 10 cm per millivolt. The ECG was recorded whenever changes occurred during or at the end of a period of hypoxemia and exercise.

Twenty one of the 43 surviving patients had a normal preoperative rest ECG.

| Preoperative stress ECG test | | | | | Postoperative stress ECG test | | | | | Clinical improvement |
|------------------------------|-----------------------------------|------------|--------------------|----------------|-------------------------------|-----------------------------------|------------|--------------------|----------------|----------------------|
| Hypoxemia | | Exercise | | | Hypoxemia | | Exercise | | | |
| Pos or neg | Max hypoxemia (% SaO_2) | Pos or neg | Max work (kpm/min) | Max heart rate | Pos or neg | Max hypoxemia (% SaO_2) | Pos or neg | Max work (kpm/min) | Max heart rate | |
| — | — | + | 100 | 107 | — | — | + | 100 | 108 | Good |
| — | 67 | + | 750 | 141 | + | 75 | — | 800 | 143 | Excellent |
| + | 78 | + | 200 | 115 | + | 78 | + | 400 | 125 | None |
| + | 77 | + | 200 | 94 | + | 84 | + | 300 | 115 | Good |
| + | 80 | + | 150 | 100 | — | 90 | + | 150 | 105 | Good |
| + | 84 | + | 200 | 115 | + | 79 | + | 400 | 120 | Good |
| + | 90 | + | 200 | 100 | + | 85 | + | 200 | 100 | Good |
| + | 73 | + | 200 | 97 | — | 78 | + | 400 | 107 | Good |
| — | 76 | + | 1 000 | 150 | — | 80 | + | 400 | 125 | Good |
| + | 73 | + | 200 | 105 | + | 67 | + | 400 | 135 | Fair |
| + | 70 | + | 400 | 100 | + | 74 | + | 400 | 105 | Good |
| + | 78 | + | 200 | 100 | + | 78 | — | 200 | 100 | Good |
| + | 84 | + | 250 | 117 | + | 88 | + | 150 | 100 | Good |
| + | 75 | + | 300 | 110 | + | 79 | — | 400 | 115 | Good |
| — | 73 | + | 500 | 120 | — | 84 | + | 300 | 120 | Excellent |
| + | 79 | + | 200 | 100 | + | 74 | + | 600 | 143 | Fair |
| + | 73 | + | 400 | 125 | + | 79 | + | 400 | 145 | Excellent |
| + | 72 | + | 200 | 107 | — | 74 | + | 400 | 125 | Good |
| + | 77 | — | 314 | 111 | — | 79 | — | 372 | 119 | Excellent |
| — | 1 6 | — | 56 | 4 | — | 1 4 | — | 45 | 3 7 | NS |
| — | 75 | — | 600 | 111 | + | 74 | — | 800 | 130 | Good |
| — | 78 | — | 1 000 | 138 | + | 70 | + | 800 | 120 | Good |
| — | 75 | — | 400 | 100 | — | 78 | + | 600 | 150 | Good |

myocardial infarct (MI) = (myocardial infarct) ICD = (intermittent claudication) delay C.B. = (collateral) Occl = (occlusion)

(Tables II and III) 18 had records showing prior myocardial infarction but isoelectric ST segments 3 had segmental ST depression in the resting ECG and 3 had intraventricular conduction delays but isoelectric ST segments.

The stress test was considered positive if either or both hypoxemia and exercise produced significant segmental ST depression (hypoxemia ≥ 0.5 mm exercise ≥ 1.0 mm). Operative ST T alterations usually disappeared by the time of the postoperative study. When the resting ECG exhibited ST depression the test was con-

sidered positive if there was additional ST segment depression of at least 1.0 mm. It is noteworthy that in all instances of positive stress tests so defined the patients experienced typical angina pectoris.

Grouping for analysis The entire group of 45 patients surviving surgery was analyzed with respect to the symptomatic result, patency of the saphenous vein graft(s) and pre and postoperative stress electrocardiography and coronary arteriography. In addition the group was subdivided according to the stress electrocardiographic result in Group I (14 pa-

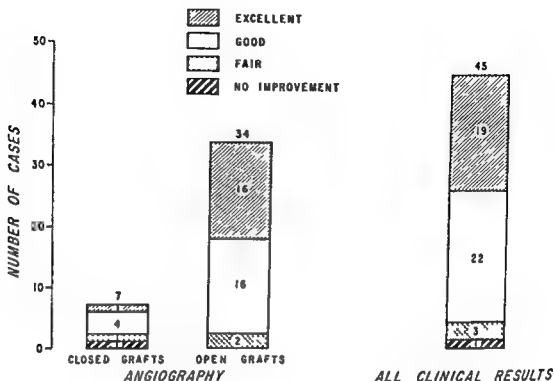


Fig. 1 Correlation of the clinical and angiographic results of aortocoronary saphenous vein bypass graft

tients) the stress ECG was positive preoperatively and became negative postoperatively in *Group II* (10 patients) both the pre and postoperative stress ECG were negative in *Group III* (15 patients) the stress ECG was positive preoperatively and remained so after surgery in *Group I* (3 patients) the stress ECG was negative preoperatively and became positive after surgery. These subgroups were examined with respect to graft patency, the distribution and evolution of obstructing coronary disease, and the maximum work and heart rates achieved before and after surgery. Each patient served as his own control. The significance of results was analyzed using the Student's *t* test.

Results

Clinical results. Fig. 1 shows the clinical results of all aortocoronary vein bypass operations. Of the 34 patients with one or more documented open grafts, 16 had an excellent result, 16 a good result, and two a fair result. In this group, 32 (94 per cent) had significant symptomatic improvement.

In one of the seven patients with closed grafts there was an excellent symptomatic result, in four a good result, in one a fair result, and in one no improvement. Thus, five of seven patients had significant clinical

improvement even though the saphenous bypass grafts were closed.

Of the entire group of 45 patients (including four not having postoperative angiography), 19 had excellent results, 22 had good results; there were three fair results, and one had no improvement. Therefore, 41 patients (91 per cent) had excellent or good symptomatic improvement.

Angiography. Angiographic findings are given in Tables II and III. As a whole, the group of 45 had significant occlusive disease of two or more coronary arteries in 31 instances. In 14 cases all significant stenoses were bypassed; in 31 cases residual stenoses could not be bypassed.

In *Group I* (stress ECG positive preoperatively, negative postoperatively) there was significant coronary disease (i.e., over 50 per cent stenosis) in at least two vessels in 11 of the 14 cases; thirteen of the 14 had double bypass grafts, and significant residual disease in another vessel was not bypassed in four instances. In *Group II* (stress ECG negative before and after surgery) seven of the 10 had significant disease in two or more coronary arteries; there were seven single grafts, and in four instances significant narrowing of other vessels could not be bypassed. Of the 18 patients in *Group III* (stress ECG positive

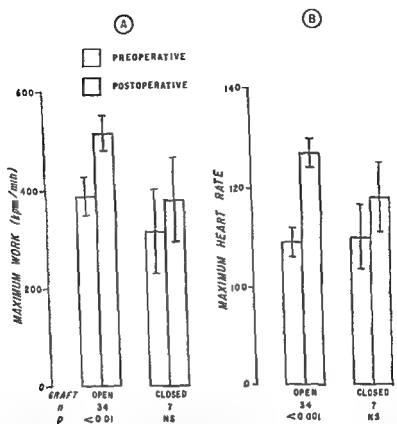


Fig 2 A and B Correlation of aortocoronary bypass graft patency with maximum work (A) and heart rate (B) achieved before and after aortocoronary bypass (the bars delimit one standard deviation of the mean)

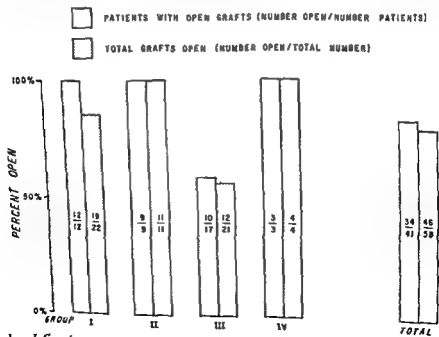


Fig 3 Correlation of aortocoronary bypass graft patency with stress ECG result sub groups

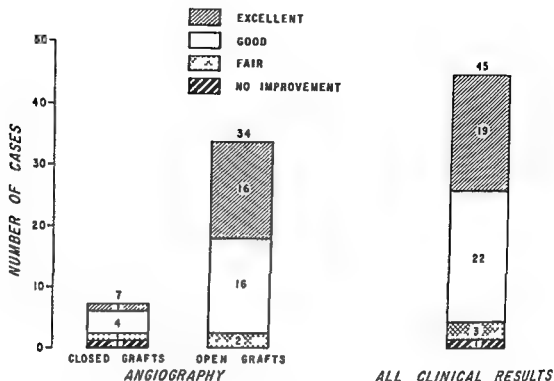


Fig. 1 Correlation of the clinical and angiographic results of aortocoronary saphenous vein bypass grafts

tients) the stress ECG was positive preoperatively and became negative postoperatively in Group II (10 patients) both the pre and postoperative stress ECG were negative in Group III (18 patients) the stress ECG was positive preoperatively and remained so after surgery in Group II (3 patients) the stress ECG was negative preoperatively and became positive after surgery. These subgroups were examined with respect to graft patency, the distribution and evolution of obstructing coronary disease and the maximum work and heart rates achieved before and after surgery. Each patient served as his own control. The significance of results was analyzed using the Student's *t* test.

Results

Clinical results Fig. 1 shows the clinical results of all aortocoronary vein bypass operations. Of the 34 patients with one or more documented open grafts, 16 had an excellent result, 16 a good result, and two a fair result. In this group, 32 (94 per cent) had significant symptomatic improvement.

In one of the seven patients with closed grafts there was an excellent symptomatic result, in four a good result, in one a fair result, and in one no improvement. Thus, five of seven patients had significant clinical

improvement even though the saphenous bypass grafts were closed.

Of the entire group of 45 patients (including four not having postoperative angiography), 19 had excellent results, 22 had good results, there were three fair results, and one had no improvement. Therefore, 41 patients (91 per cent) had excellent or good symptomatic improvement.

Angiography Angiographic findings are given in Tables II and III. As a whole, the group of 45 had significant occlusive disease of two or more coronary arteries in 31 instances. In 14 cases, all significant stenoses were bypassed; in 31 cases, residual stenoses could not be bypassed.

In Group I (stress ECG positive preoperatively, negative postoperatively) there was significant coronary disease (i.e., over 50 per cent stenosis) in at least two vessels in 11 of the 14 cases; thirteen of the 14 had double bypass grafts and significant residual disease in another vessel was not bypassed in four instances. In Group II (stress ECG negative before and after surgery), seven of the 10 had significant disease in two or more coronary arteries; there were seven single grafts and in four instances significant narrowing of other vessels could not be bypassed. Of the 18 patients in Group III (stress ECG positive

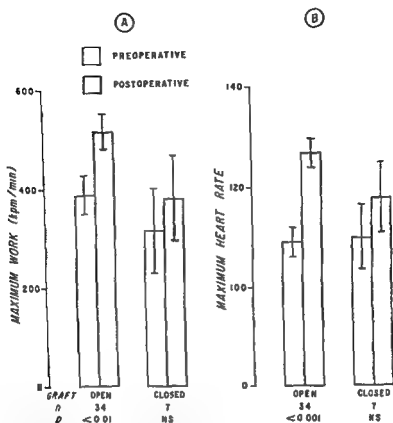


Fig 2 A and B Correlation of aortocoronary bypass graft patency with maximum work (A) and heart rate (B) achieved before and after aortocoronary bypass (the bars delimit one standard deviation of the mean)

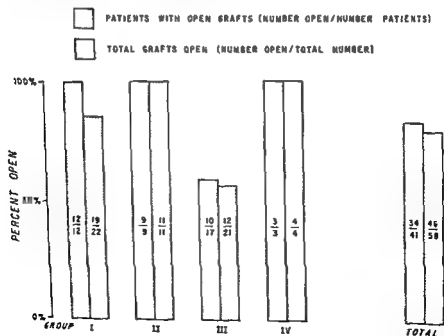


Fig 3 Correlation of aortocoronary bypass graft patency with stress ECG result subgroups

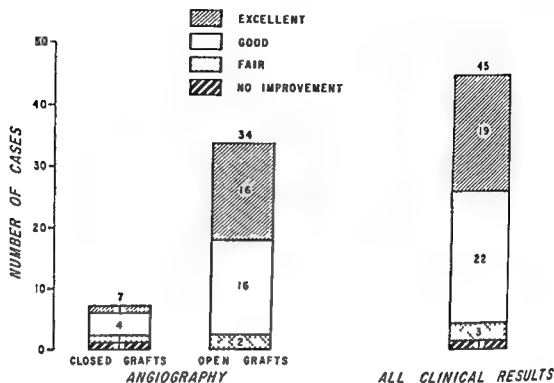


Fig. 1 Correlation of the clinical and angiographic results of aortocoronary saphenous vein bypass graft

tients) the stress ECG was positive preoperatively and became negative postoperatively in Group II (10 patients) both the pre and postoperative stress ECG were negative in Group III (18 patients) the stress LCC was positive preoperatively and remained so after surgery in Group II (3 patients) the stress LCC was negative preoperatively and became positive after surgery. These subgroups were examined with respect to graft patency, the distribution and evolution of obstructing coronary disease, and the maximum work and heart rates achieved before and after surgery. Each patient served as his own control. The significance of results was analyzed using the Student *t* test.

Results

Clinical results Fig. 1 shows the clinical results of all aortocoronary vein bypass operations. Of the 34 patients with one or more documented open grafts 16 had an excellent result, 16 a good result, and two a fair result. In this group, 32 (94 per cent) had significant symptomatic improvement.

In one of the seven patients with closed grafts there was an excellent symptomatic result, in four a good result, in one a fair result, and in one no improvement. Thus five of seven patients had significant clinical

improvement even though the saphenous bypass grafts were closed.

Of the entire group of 45 patients (including four not having postoperative angiography) 19 had excellent results, 22 had good results, there were three fair results, and one had no improvement. Therefore 41 patients (91 per cent) had excellent or good symptomatic improvement.

Angiography Angiographic findings are given in Tables II and III. As a whole the group of 45 had significant occlusive disease of two or more coronary arteries in 31 instances. In 14 cases all significant stenoses were bypassed; in 31 cases residual stenosis could not be bypassed.

In Group I (stress ECG positive preoperatively, negative postoperatively) there was significant coronary disease (i.e. over 50 per cent stenosis) in at least two vessels in 11 of the 14 cases; thirteen of the 14 had double bypass grafts, and significant residual disease in another vessel was not bypassed in four instances. In Group II (stress LCC negative before and after surgery) seven of the 10 had significant disease in two or more coronary arteries; there were seven single grafts, and in four instances significant narrowing of other vessels could not be bypassed. Of the 18 patients in Group III (stress ECG positive

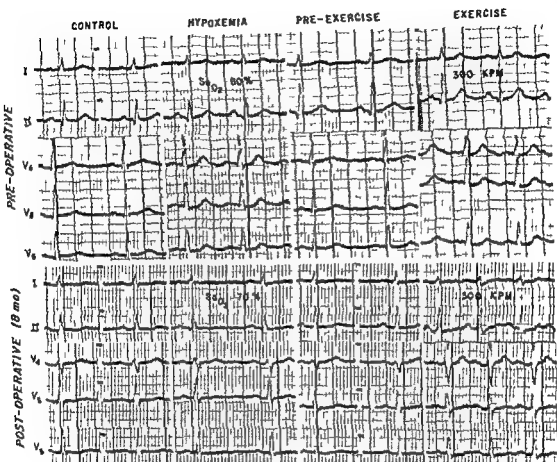


Fig 5 Stress ECG of patient 5 (A-D) Group I. The preoperative ECG (upper panel) shows segmental ST depression most marked in V_1 and V_2 with hypoxemia. Postoperatively (lower panel) there is no significant segmental ST depression even at greater hypoxemia.

imum work was 379 ± 87 kpm per minute (21 per cent increase $p = NS$). The mean maximum heart rate achieved by this group preoperatively was 110 ± 6 compared to the postoperative mean maximum rate of 118 ± 7 (7 per cent increase $p = NS$) (Fig 2).

Of the 34 patients with one or more grafts open (Table I, Fig 1) 22 had positive stress ECG tests before surgery and 13 were positive afterwards (59 per cent). The preoperative mean maximum work performed by this group was 398 ± 38 kpm per minute and the postoperative mean maximum work was 515 ± 35 kpm per minute (33 per cent increase $p < 0.01$). The mean maximum heart rate achieved preoperatively was 109 ± 7 compared to the postoperative mean maximum rate of

127 ± 3 (17 per cent increase $p < 0.001$) (Fig 2).

All patients were stressed by the same degree of hypoxemia both pre and postoperatively.

In general, when one stress test was positive, so was the other. In about 10 per cent of instances there was a dichotomy: preoperatively the hypoxemia test was positive four times when the exercise test was negative and the exercise test was positive four times when the hypoxemia test was negative; postoperatively the hypoxemia test was positive four times when the exercise test was negative and the exercise test was positive six times when the hypoxemia test was negative.

The stress electrocardiographic results are more meaningful when the entire group

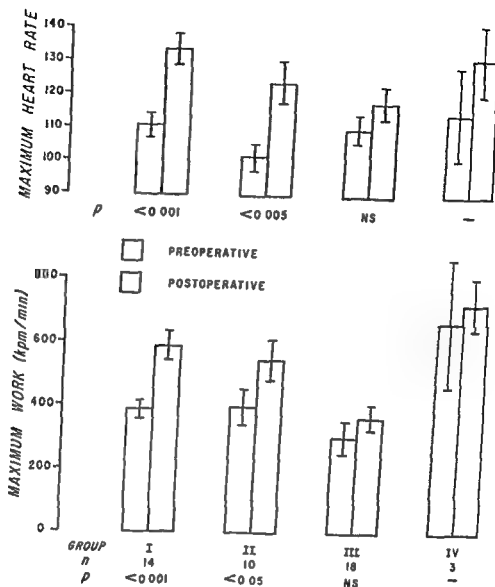


Fig. 4 Relationship between stress ECG result subgroup and maximum work (lower panel) and heart rate (upper panel) achieved before and after aortocoronary bypass (the bars delimit one standard deviation of the mean)

before and after surgery) significant disease of two or more major coronary arteries was present in 10 cases. 15 of the 18 had single bypass grafts and significant residual disease of other vessels was not bypassed in 10 instances. Of the three patients in Group IV (stress ECG negative preoperatively, positive postoperatively) two had single grafts and one had double grafts; in all cases there was significant disease of at least two vessels and residual stenosis was not bypassed in all instances.

Postoperative angiography was done in 41 patients between one and 12 months after surgery, averaging five months (Tables I, II, and III). One or more grafts were open in 34 (83 per cent) and seven patients had no open graft. Of all the saphenous

bypass grafts, 46 of 56 (79 per cent) were patent (Fig. 3).

Eight patients had interval occlusion or progression of proximal stenosis in the grafted coronary artery. Five had interval occlusion of other non-grafted vessels and one developed significant intercoronary collaterals after surgery. In three instances there was more than 50 per cent stenosis in the coronary artery distal to the bypass graft.

Stress electrocardiography The seven patients with closed grafts (Table I, Fig. 1) had positive stress ECG results both before and after surgery. The preoperative mean maximum work performed was 314 ± 84 (standard error of the mean) kpm per minute and the postoperative mean max

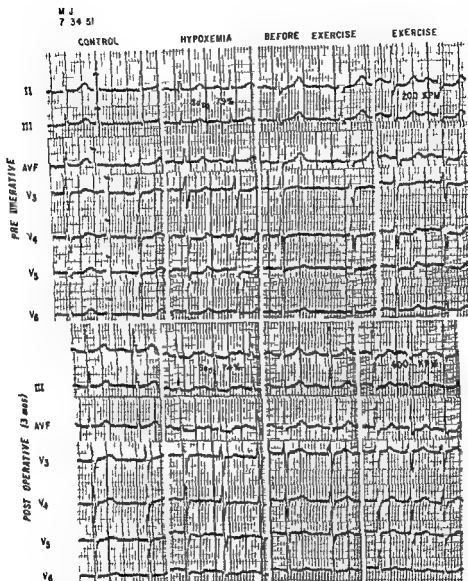


Fig. 7. Stress ECG of patient 16 (M J) Group III. The preoperative hypoxemia and exercise tests (upper panel) are highly positive. Postoperatively (lower panel) the hypoxemia test is equally positive, the exercise test less so, both at greater level of hypoxemia and exercise.

and 11 (53 per cent) had objective evidence of improvement in terms of negative post-operative stress electrocardiograms and the achievement of significantly more work and greater heart rates after surgery. All of these patients, who had preoperative angiography (21) were found to have at least one open type I graft (Groups III and IV) (47 per cent) were objectively either no better or were worse after surgery, had positive postoperative stress electrocardio-

grams and were unable to achieve significantly greater work or heart rates after surgery. Only 13 of the 20 patients (65 per cent) in these groups had patent grafts.

Discussion

The evaluation of any myocardial revascularization technique is complicated by the symptomatic variability of the disease and by the influence of a number of interacting variables which tend to obscure



Fig 6 A through C Coronary angiograms of patient whose stress ECG is depicted in Fig 5. There is occlusion of the left anterior descending coronary (A preoperative and B postoperative) successfully bypassed in C.

of 45 patients is subdivided on the basis of the preoperative test result and the change (if any) which occurred after surgery. In Group I the stress ECG was positive before surgery and became negative postoperatively in this group of 14 patients (31 per cent). Twelve of these patients had postoperative angiograms showing patent grafts in all cases. There was a total of 22 grafts in these 14 patients and 19 grafts were open (Fig 3). Most patients in Group I were able to perform significantly more work and achieve significantly greater heart rates postoperatively (Fig 4). Preoperatively the group accomplished a mean maximum work level of 389 ± 32 kpm per minute and postoperatively 559 ± 43 kpm per minute (51 per cent increase, $p < 0.001$). Before surgery they achieved a mean maximum heart rate of 111 ± 4 after surgery 134 ± 5 (21 per cent increase, $p < 0.001$). Figs 5 and 6 illustrate pre and postoperative angiograms and stress electrocardiograms in a patient of Group I.

In Group II the stress ECG was negative both before and after surgery (10 patients, 22 per cent). Postoperative angiography in nine of these patients showed that all bypass grafts were patent (Fig 3). Most of the patients in this group performed significantly more work and reached a significantly greater heart rate postoperatively (Fig 4). Preoperatively the group reached a mean maximum work level of 400 ± 57 kpm per minute postoperatively $470 \pm$

63 kpm per minute (43 per cent increase, $p < 0.05$). The mean maximum heart rate increased from 102 ± 4 preoperatively to 125 ± 6 postoperatively (23 per cent increase, $p < 0.005$).

The 18 patients in Group III (40 per cent of the total) had positive stress electrocardiograms both before and after surgery. Seventeen of these had postoperative angiography, showing that 10 patients had one or more grafts open and that 12 of the 21 total grafts in this group were open (Fig 3). Most patients in this group were not able to generate significantly more work or greater heart rate after surgery (Fig 4). Preoperatively the group reached a mean maximum work level of 314 ± 56 kpm per minute and postoperatively 372 ± 43 kpm per minute (18 per cent increase, $p = \text{NS}$). The pre and postoperative maximum heart rates achieved were 111 ± 4 and 119 ± 4 respectively (7 per cent increase, $p = \text{NS}$). Figs 7 and 8 illustrate pre and postoperative angiograms and stress electrocardiograms in a patient of Group III.

Group IV comprised three of the 45 patients (7 per cent) whose stress ECG was negative preoperatively and positive after surgery. All three had open bypass grafts (Fig 3). As a group they achieved higher work levels and heart rates both before and after surgery (Fig 4) but further comparisons with the other groups are invalided by the small number.

In summary, the 24 patients in Groups I

residual non bypassed coronary obstruction

Patients in Group II with postoperative negative stress ECG tests and increased heart rates and work production illustrated the need to quantitate the work and heart rates achieved during exercise since some patients with bona fide coronary disease had negative stress electrocardiography or the tests were aborted by chest pain before discriminating ECG changes occurred. Although most of the patients in this group had single grafts there was a relatively low incidence of residual non bypassed coronary disease and like Group I all patients had one or more patent grafts. Taken together Groups I and II showed good correlation between negative postoperative stress ECG tests and improved coronary circulation resulting from at least one patent aortocoronary graft.

The patients in Groups III and IV had positive postoperative stress ECG tests and collectively were unable to generate significantly greater heart rates or work in comparison to the levels achieved preoperatively. The persistently abnormal stress results in Group III correlated with the relatively low incidence of open grafts (6 per cent) and the larger number of coronary obstructions which were not bypassed. Group IV patients reversed their originally negative stress ECG tests becoming positive postoperatively although they were able to do more work and achieve greater heart rates than before surgery. All of the patients in Group IV had significant disease of other vessels and two of the three had postoperative coronary occlusions.

It seems clear from these simple and non-invasive studies that stress electrocardiography and quantitation of the heart rate and work achieved by exercise provide a relatively accurate objective assessment of the results of aortocoronary bypass surgery. Although over 90 per cent of patients were improved symptomatically and nearly 80 per cent of the vein bypass grafts were patent at the time of postoperative angiography, only 53 per cent were objectively improved.

The predictive value of stress electrocardiography was especially high. The graft patency rate was 90 to 100 per cent

when (1) the stress ECG was positive preoperatively and reversed postoperatively or (2) when the stress ECG was negative both before and after surgery and the patients could achieve significantly greater work and heart rates postoperatively. On the other hand when the stress test remained positive or became so after surgery there was only about 65 per cent probability that one or more grafts were open and a high probability of significant residual coronary obstruction. Moreover in all instances in which there was no open graft after surgery the postoperative stress ECG was positive and there was no significant increase in heart rate and work over that achieved by preoperative exercise.

Summary

Forty-five patients having saphenous vein aortocoronary bypass surgery were studied prospectively by pre and postoperative hypoxemia and graded exercise electrocardiography. The stress electrocardiographic results were correlated with the degree of symptomatic relief and the angiographic evidence of graft patency and the distribution and evolution of coronary occlusive disease.

Forty-one of the total group of 45 patients (91 per cent) and 32 of 34 (94 per cent) with one or more documented open grafts had excellent or good symptomatic improvement. Incongruously five of seven patients with closed grafts also had significant clinical improvement.

Functional improvement was documented in 53 per cent of patients after aortocoronary bypass surgery in terms of postoperative reversal of the preoperatively positive stress ECG or persistence of a negative stress ECG together with the achievement of significantly greater mean maximum exercise and heart rate. All of these patients had one or more patent saphenous vein bypass grafts. Forty-seven per cent of the patients failed to display objective evidence of improvement postoperatively, the stress ECG either remained positive or became so and collectively the group was unable to significantly increase the mean maximum exercise level or heart rate in comparison to that achieved before surgery. Only 65 per cent of these



Fig 8 1 through D Coronary angiograms of patient whose stress ECG is depicted in Fig 7 The right coronary is normal (1) and there is 90 per cent stenosis of the left anterior descending (B) successfully bypassed in (C) and (D)

whether myocardial ischemia is in fact diminished after surgery. Indeed pain relief after operation may be at least in some part the result of the period of enforced bed rest during pre and postoperative circulatory infarction of a pain producing zone of ischemic myocardium and postoperative physical and emotional conditioning. Although postoperative angiography verifies the patency of saphenous vein aortocoronary bypass grafts it provides inadequate quantitation of the degree of functional improvement which results from the balance between increased coronary flow provided by the bypass graft and residual coronary obstruction which could not be bypassed. Moreover it may not be feasible or desirable to do postoperative angiography in all cases. For these reasons standardized hypoxemia and exercise electrocardiography was employed as an objective measurement of the functional result of aortocoronary bypass surgery. The reproducibility of hypoxemia¹⁹ and exercise^{21, 22} electrocardiography has been established as long as exercise is graded from levels below the threshold for a positive response is maintained for at least three minute periods, and the patients are not undergoing physical training.³

The majority of patients in this study had significant symptomatic improvement. Over 90 per cent had excellent or good clinical results, and about 80 per cent of all grafts were patent when examined angiographically five or six months after surgery. These results are quite similar to those reported by others.^{21, 22} Incongruously, most

of the patients with occluded grafts also were improved symptomatically after surgery.

Stress electrocardiography provided more discriminating evaluation of the functional results of surgery. The patients with closed aortocoronary bypass grafts exhibited persistently positive stress electrocardiography and as a group failed to achieve significantly greater heart rates and exercise when tested postoperatively. On the other hand the mean maximum heart rate and work achieved as a group by those with one or more grafts open increased significantly after surgery, although stress electrocardiography was positive in almost half of the patients after surgery. Normalization of the exercise ECG and increase in the heart rate-blood pressure product have also been observed by Amsterdam and colleagues²⁴ and by Bergman and associates²⁵ in the majority of patients examined early after aortocoronary saphenous vein bypass.

In this study the complicated interrelationship between saphenous vein bypass graft patency and residual coronary obstruction was best expressed in relation to the difference if any between preoperative and postoperative stress electrocardiography. The patients in Group I reversed their originally positive preoperative stress ECG becoming negative after surgery and generally achieved significantly greater heart rates and exercise postoperatively. Although all of the patients had one or more grafts open this group also had the greatest number of double bypass grafts and the least

- reproducibility and follow up results. *Circulation* 43 and 44 (Suppl 11) 16 1971
- 23 Ledwood D R, Rosing D R and Epstein S F. Circulatory and symptomatic effects of physical training in patients with coronary artery disease and angina pectoris. *N Engl J Med* 286 939 1972
- 24 Amsterdam E A, Iben A, Hurley B J, Mansour E, Hughes J L, Salel A F, Zelis R and Mason D T. Saphenous vein bypass graft for refractory angina pectoris: physiologic evidence for enhanced blood flow to the ischemic myocardium. *Am J Cardiol* 26 623 1970
- 25 Bergman S A, Urschel H C Jr and Blomqvist G. Pre and postoperative exercise testing in patients undergoing direct myocardial revascularization. *Circulation* 43 and 44 (Suppl 11) 141 1971

subjects had one or more patent bypass grafts

Stress electrocardiography and quantitation of the maximum work and heart rate achieved by exercise provide accurate assessments of the relief of myocardial ischemia produced by myocardial revascularization. In this study, when the postoperative stress ECG was negative and the patient was able to achieve significantly greater exercise and heart rate the probability of patency of one or more bypass grafts was virtually certain. On the other hand, when the postoperative stress ECG was positive and little or no increase in exercise and heart rate was possible there was high probability of graft occlusion or significant residual coronary disease.

The authors are grateful to their surgical colleagues Drs. Albert Starr, Robert P. Hodum, Lawrence I. Bonchek, and Richard P. Anderson. Mr. Larry Willett and Mrs. Barbara Burk provided invaluable assistance with electrocardiography and data analysis.

REFERENCES

- 1 Favoloro R G Saphenous vein graft in the surgical treatment of coronary artery disease operative technique *J Thorac Cardiovasc Surg* 50:178 1969
- 2 Favoloro R G Effler D B Groves L H Sheldon W C Shires F A and Sones F M Jr Severe segmental obstruction of the left main coronary artery and its divisions surgical treatment by saphenous vein graft technique *J Thorac Cardiovasc Surg* 60:469 1970
- 3 Favoloro R G Effler D B Groves L H Sheldon W C and Sones F M Jr Direct myocardial revascularization by saphenous vein graft present operative technique and indications *Ann Thorac Surg* 10:97 1970
- 4 Johnson W D Flemmer R J Lepley D Jr and Ellison E H Extended treatment of severe coronary artery disease a total surgical approach *Ann Surg* 170:160 1969
- 5 Johnson W D and Lepley D Jr An aggressive surgical approach to coronary artery disease *J Thorac Cardiovasc Surg* 59:147 1970
- 6 Mitchell H F Adam M Lambert C J Sungu U and Shiekh S Ascending aorta to coronary artery saphenous vein bypass grafts *J Thorac Cardiovasc Surg* 60:457 1970
- 7 Anderson R P Hodum R Wood J and Starr A Direct revascularization of the heart early clinical experience with 200 patients *J Thorac Cardiovasc Surg* 63:353 1972
- 8 Morris G C Reul G J Howell J F Crawford E S Chapman D W Beazley H L Winters W L Peterson P K and Lewis J M Follow up results of distal coronary artery bypass for ischemic heart disease *Am J Cardiol* 29:180 1972
- 9 Reul G J Morris G C Howell J F Crawford E S Sandford J M and Wukitch D C Experience with coronary artery bypass grafts in the treatment of coronary artery disease *Surgery* 71:586 1972
- 10 Dimelzon G K Gau G T and Davis G D Early results of vein bypass grafts for coronary artery disease *Circulation* 43 and 44 (Suppl II) 101 1971
- 11 Grondin C M Lepige G Castonguay Y R Meere C and Grondin P Aorto-coronary bypass graft initial blood flow through the graft and early postoperative patency *Circulation* 44:815 1971
- 12 Urschel H C Solis R M Miller E R Rizzuk M A and Wood R E Factors influencing flow through aortocoronary artery sphenous vein bypass grafts *Am J Cardiol* 29:295 1972
- 13 Walker J A Friedberg H D Flemmer R J and Johnson W D Determinants of angiographic patency of aortocoronary vein bypass grafts *Circulation* 45 and 46 (Suppl I) 86 1971
- 14 Johnson W D Flemmer R J Manley J C and Lepley D Jr The physiologic parameters of ventricular function as affected by direct coronary surgery *J Thorac Cardiovasc Surg* 60:183 1970
- 15 Camperio L Alonzo F Ellis G and Bourassa G Left ventricular performance during exercise before and after aorto-coronary vein graft surgery *Circulation* 43 and 44 (Suppl II) 148 1971
- 16 Rees G Bristow J D Kreimkau E L Green G S Herr R H Griswold H E and Starr A Influence of aortocoronary bypass surgery on left ventricular performance *N Engl J Med* 284:1116 1971
- 17 Manley J C Johnson W D Flemmer R J and Lepley D Objective evaluation of the effects of direct myocardial revascularization on ventricular performance utilizing submaximal ergometer exercise testing *Am J Cardiol* 29:277 1972
- 18 Chatterjee K Swan H J C Parmley W W Sustutis H Marcus H and Mitloff J Depression of left ventricular function due to acute myocardial ischemia and its reversal after aortocoronary sphenous vein bypass *N Engl J Med* 286:1117 1972
- 19 Kassebaum D G Judkins M P and Griswold H E Stress electrocardiography in the evaluation of surgical revascularization of the heart *Circulation* 40:297 1969
- 20 Judkins M P Selective coronary arteriography a percutaneous transluminal technique *Radiology* 89:815 1967
- 21 Redwood D R Roseng D R Goldstein R F Besser G D and Epstein S L Importance of the design of an exercise protocol in the evaluation of patients with angina pectoris *Circulation* 43:618 1971
- 22 Blomqvist G and Atkins J M Repeated exercise testing in patients with angina pectoris

in identifying slurs in the waveform but that the identification of a notch was much less subjective and was consistently identified by several observers. Therefore we decided to re-examine our initial group of subjects to determine whether appropriate new diagnostic criteria could be developed. We were successful in developing satisfactory new criteria. 966 subjects who received routine health examinations at the University of Pennsylvania Health Evaluation Center were studied by wide band recording of the electrocardiogram using our new criteria.

Material and methods

1 Original subjects Two groups which had been studied prior to 1966 were selected for further analysis. Group A consisted of the 100 normal controls¹ who were healthy male employees of the Provident Mutual Life Insurance Company. This group received annual health maintenance examinations. High frequency records from these subjects have been obtained at intervals of two to three years since 1950 and were followed up to 1966. records made prior to 1956 were still available and were used in this study. The ages at entry were 38 to 63 with a mean age of 51 years. Myocardial infarctions² occurred in three members of this group in the interval 1955 to 1966; all the others have remained free from heart disease. Since 1966 24 new male subjects were added to our group. Two of the new entrants developed coronary artery disease with myocardial infarction; their records will be discussed separately.

Group B consisted of 16 ambulatory subjects who had recovered very satisfactorily from a well documented myocardial infarction³; their ages were 43 to 73 years with a mean age of 56 years. Six of these subjects were female.

2 Health Evaluation Center studies Since 1966 a series of high frequency electrocardiograms have been obtained on 966 ambulatory male subjects at the Health Evaluation Center at the Hospital of the University of Pennsylvania. These patients were studied by the usual clinical methods including chest x-ray, conventional electrocardiogram, blood chemistry, hemoglobin, white blood count, and other non-invasive techniques as deemed appropriate. The

ages were 35 to 72 years with a mean age of 57 years. Eight hundred seventy five (875) of these subjects were classified as normal. 24 had documented histories of myocardial infarction. 12 had a diagnosis of angina pectoris. 35 had systolic blood pressure greater than 160 mm Hg and diastolic blood pressure higher than 100 mm Hg. 20 had a cholesterol level of 300 mg per cent or higher.

Methods

Records from Group A (100 subjects) plus Group B (76 subjects) were made by photographing the face of a cathode ray oscilloscope on 1 inch wide rapidly moving (350 mm per second) photographic paper.⁴ Records from the Health Evaluation Center group (966 people) plus 24 new subjects from the insurance company were made by photographing the face of a cathode ray oscilloscope with continuously moving 16 mm film.⁵ Examples of high frequency notching, utilizing this technique are shown in Figs 1 and 2. Excluding major changes of directions of QRS such as peaks of the R and nadirs of the Q and S waves, slurs and notches are defined as changes of the slope of QRS. Slurs are departures from the smooth slope without change of sign, whereas notches are changes both of slope and sign. Since 1969 records have been made with a Linac II computer. The output of this computer is displayed on an oscilloscope and three consecutive complexes are photographed with a polaroid camera.⁶ Comparison of records obtained using both the moving film and the computer has indicated that the configuration of the waveform is essentially identical. The frequency response of all three systems exceeded a range flat from 0.15 to 1000 Hz. Standard scalar limb and precordial leads were used in all wide band studies.

Results

Analysis of original 176 subjects (Groups A and B) All notches and slurs in the three largest limb leads and the six precordial leads in these 176 subjects are tabulated in Table I. Table II is based on identical data but omits all information from Leads V₁, V₂, and V₃. Notches only appearing in the three largest limb leads and in V₄, V₅, and V₆ are tabulated in Table III. It can

Wide band recording of the electrocardiogram and coronary heart disease

Paul H Langner Jr MD, IACP

David B Geselowitz PhD*

Stanley A Briller MD

Philadelphia Pa

Wide band recording of the electrocardiogram utilizing an expanded time scale and greater amplitude^{1,2} than the conventional method is a useful tool because it increases the detection rate of myocardial disease. Our previous studies have demonstrated that one of the chief values of this method is its unique capability of recording high frequency notches and slurs which are much more common in the QRS complex of patients with clinical and preclinical coronary heart disease than in apparently healthy subjects.³⁻⁶ These high frequency notches and slurs are not revealed by the conventional electrocardiogram even when its frequency response is raised to 100 Hz. Other investigators have published studies utilizing high frequency electrocardiography which have confirmed these findings of increased notching in heart disease. This work includes studies of patients with coronary artery disease^{10,12} patients with cardiomyopathies¹⁸ biventricular hypertrophy¹⁷ and rheumatic heart disease.¹⁹ Of these investigators Burch and co workers^{10,11} and Selvester and co workers¹² used the spatial vectorcardio-

gram. Franke and Braunstein¹⁸ used power spectrum analysis. Flowers, Horan and colleagues^{19,17} made extensive studies using three orthogonal leads in both coronary heart disease and biventricular hypertrophy. Reynolds and co workers¹⁵ employed a modified spatial vectorcardiogram.

Our earlier studies indicated that people without heart disease could be distinguished from subjects with coronary heart disease by enumerating the total number of notches and slurs in the three largest limb leads and the six precordial V leads.⁹ A similar method was initiated in a program involving persons seen at the Health Evaluation Center at the Hospital of the University of Pennsylvania to expand these studies to a large group of subjects. As this study progressed we found a considerable overlap in the notch plus slur count in V₁, V₂ and V₃ between the so called normal group which by non-invasive clinical techniques revealed no evidence of coronary heart disease and a group of subjects having well documented coronary heart disease. It had also become apparent to us that there was some ambiguity

From the Cardiovascular Pulmonary Division, Section of Cardiology, Department of Medicine, University of Pennsylvania School of Medicine, Philadelphia, Pa.

Supported by an award from the United States Public Health Service Grant HL 10094.

Received for publication Oct. 30, 1972.

Reprint requests to Dr. Stanley A. Briller, 937 Gates Bldg., West Hospital of the University of Pennsylvania, 3400 Spruce St., Philadelphia, Pa. 19104.

*Present address: Department of Bioengineering, Pennsylvania State University, University Park, Pa.

September 1973 Vol. 85 No. 3 pp. 308-317

1 of m 86
N of bc 3

Table I Combined number high frequency notches and slurs in nine leads

| Notches and slurs | Normal subjects | Abnormal subjects | Notches and slurs | Normal subjects | Abnormal subjects |
|-------------------|-----------------|-------------------|-------------------|-----------------|-------------------|
| 0 | 2 | 0 | 21 | 0 | 4 |
| 1 | 0 | 0 | 22 | 0 | 10 |
| 2 | 2 | 0 | 23 | 0 | 2 |
| 3 | 4 | 0 | 24 | 0 | 3 |
| 4 | 5 | 0 | 25 | 0 | 5 |
| 5 | 11 | 0 | 26 | 0 | 2 |
| 6 | 9 | 0 | 27 | 0 | 0 |
| 7 | 9 | 0 | 28 | 0 | 2 |
| 8 | 13 | 0 | 29 | 0 | 4 |
| 9 | 13 | 0 | 30 | 0 | 1 |
| 10 | 11 | 2 | 31 | 0 | 6 |
| 11 | 9 | 1 | 32 | 0 | 4 |
| 12 | 6 | 4 | 33 | 0 | 2 |
| 13 | 2 | 4 | 34 | 0 | 0 |
| 14 | 1 | 2 | 35 | 0 | 1 |
| 15 | 1 | 2 | 36 | 0 | 0 |
| 16 | 2 | 3 | 37 | 0 | 0 |
| 17 | 0 | 0 | 38 | 0 | 0 |
| 18 | 0 | 4 | 39 | 0 | 1 |
| 19 | 0 | 1 | 40 | 0 | 0 |
| 20 | 0 | 5 | 41 | 0 | 1 |
| | | | Totals | 100 | 76 |

notches plus slurs in six leads counting notches plus slurs and six leads counting notches alone) can differentiate normals from abnormals ($P < 0.001$). For screening purposes the first procedure was chosen employing the three largest limb leads and V_1 , V_4 and V_6 counting notches only. This decision validated by the foregoing statistical analysis was justified by the fewer number of leads and events to be counted and by simpler incorporation into a computerized system to be reported later.

Health Evaluation Center Data. Results for the Health Evaluation Center subjects are summarized in Table IV. On clinical ground 387 subjects were classified as normal. Of this clinically normal group 89 per cent had three notches or less and 11 per cent had a notch count of four or more. Approximately 300 of these subjects had two or more high frequency electrocardiogram at yearly intervals for four years. None of these exhibited any significant clinical change except for one subject who developed myocardial infarction and is included in the infarction group (Fig. 3).

There were 74 patients with well documented histories of myocardial infarction

and all but two had abnormal conventional electrocardiograms. As shown in Table IV 79 per cent of the 24 subjects had an abnormal number of notches. Repeated high frequency records at intervals of a year or more showed no serial changes. There were 12 subjects with a diagnosis of angina pectoris of these seven had an abnormal number of notches. The electrocardiogram of one of these subjects is seen in Fig. 4. The first two groups are hypertension (160/100 or higher—37 subjects) and hypercholesterolemia (300 mg. per cent or higher—20 subjects). Notching in these two categories was not higher than in the normal group. All members of the group (Table IV) were active and gainfully employed. No one had a history of congestive heart failure. Only three of the 35 subjects with hypertension had mild to moderate left ventricular hypertrophy. Of the three subjects with left ventricular hypertrophy by conventional electrocardiographic standards none had excess high frequency notching.

Additional Provident Mutual Life Insurance Company employees. All 24 of these subjects had normal conventional low frequency and high frequency electro-

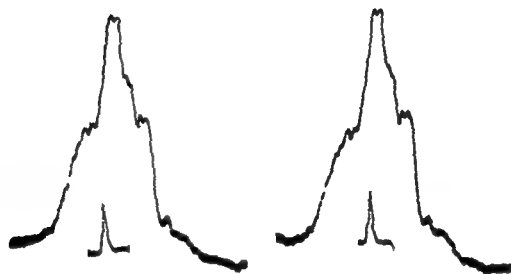


Fig 1 Two consecutive high frequency complexes (Lead II) with the conventional recording below. There is obvious notching in the large high frequency tracing which cannot be identified in the conventional records below.

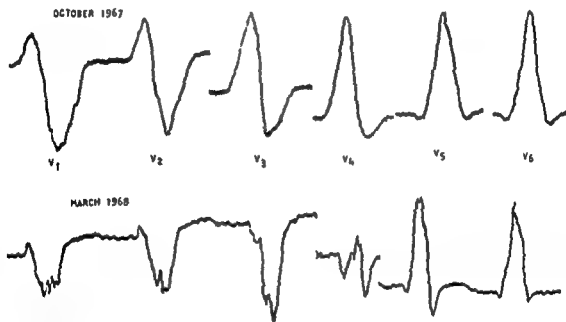


Fig 2 The upper panel consists of high frequency chest lead made in 1967 prior to infarction. The bottom panel shows high frequency records made from the same precordial sites after infarction. Lead V_2 consists of a Q S complex and V_4 shows obvious low frequency serial change. None of the notching evident in the lower tracing was present in the conventional electrocardiogram of the same date.

be seen in Table I that 94 per cent of normals have 12 or fewer notches and slurs and 91 per cent of those with heart disease have more than 12 notches and slurs. In Table II 89 per cent of subjects without heart disease have six or less notches and slurs whereas 89 per cent of those with heart disease had more than six notches and slurs. Table III contains the number of notches

(slurs omitted) in the three largest limb leads and in V_1 , V_2 , and V_6 . Ninety two per cent of those without heart disease had three notches or less but 93 per cent of subjects with heart disease had more than three notches. Results from Tables I, II, and III were examined statistically. The Median Test²² confirmed that each of the three procedures (nine leads counting

Table I Combined number high frequency notches and slurs in nine leads

| Notches and slurs | Normal subjects | Abnormal subjects | Notches and slurs | Normal subjects | Abnormal subjects |
|-------------------|-----------------|-------------------|-------------------|-----------------|-------------------|
| 0 | 2 | 0 | 21 | 0 | 4 |
| 1 | 0 | 0 | 22 | 0 | 10 |
| 2 | 2 | 0 | 23 | 0 | 2 |
| 3 | 4 | 0 | 24 | 0 | 3 |
| | 5 | 0 | 25 | 0 | 5 |
| 5 | 11 | 0 | 26 | 0 | 2 |
| 6 | 9 | 0 | 27 | 0 | 0 |
| 7 | 9 | 0 | 28 | 0 | 2 |
| 8 | 13 | 0 | 29 | 0 | 4 |
| 9 | 13 | 0 | 30 | 0 | 1 |
| 10 | 11 | 2 | 31 | 0 | 1 |
| 11 | 9 | 1 | 32 | 0 | 4 |
| 12 | 6 | 4 | 33 | 0 | 2 |
| 13 | 2 | 4 | 34 | 0 | 0 |
| 14 | 1 | 2 | 35 | 0 | 1 |
| 15 | 1 | 2 | 36 | 0 | 0 |
| 16 | 2 | 3 | 37 | 0 | 0 |
| 17 | 0 | 0 | 38 | 0 | 0 |
| 18 | 0 | 4 | 39 | 0 | 1 |
| 19 | 0 | 1 | 40 | 0 | 0 |
| 20 | 0 | 5 | 41 | 0 | 1 |
| Total | | | | 100 | 16 |

notches plus slurs in 12 leads counting notches plus slurs and six leads counting notches alone) can differentiate normals from abnormal (1 < 0.001). For screening purposes the list procedure was chosen employing the three largest limb leads and V_1 , V_4 and V_6 counting notches only. This decision validated by the foregoing statistical analysis was justified by the fewer number of leads and events to be counted and by simpler incorporation into a computerized system to be reported later.

Health Evaluation Center Data: Results for the Health Evaluation Center subjects are summarized in Table IV. On clinical ground 873 subjects were classified as normal. Of this clinically normal group 89 per cent had three notches or less and 11 per cent had a notch count of four or more. Approximately 300 of these subjects had two or more high frequency electrocardiograms at yearly intervals for four years. None of these exhibited any significant serial change except for one subject who developed myocardial infarction and is included in the infarction group (Fig. 3).

There were 24 patients with well documented myocardial infarction

and all but two had abnormal conventional electrocardiograms. As shown in Table IV 79 per cent of the 24 subjects had an abnormal number of notches. Reported high frequency records at intervals of a year or more showed no serial changes. There were 12 subjects with a diagnosis of angina pectoris of these seven had an abnormal number of notches. The electrocardiogram of one of these subjects is seen in Fig. 4. The first two groups are hypertension (160/100 or higher—33 subjects) and hypercholesterolemia (300 mg per cent or higher—20 subjects). Notching in these two categories was not higher than in the normal group. All members of the group (Table IV) were active and carefully employed. No one had a history of congestive heart failure. Only three of the 33 subjects with hypertension had mild to moderate left ventricular hypertrophy. Of the three subjects with left ventricular hypertrophy by conventional electrocardiographic standards none had excess high frequency notching.

Additional Provident Mutual Life Insurance Company employees: All 24 of these subjects had normal conventional low frequency and high frequency electro-

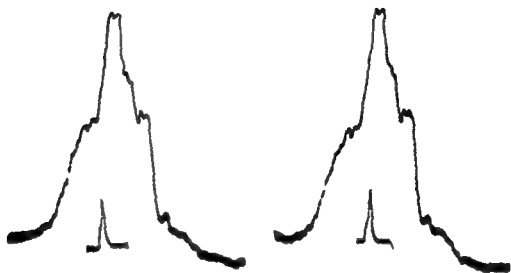


Fig 1 Two consecutive high frequency complexes (Lead II) with the conventional recordings below. There is obvious notching in the large high frequency tracing, which cannot be identified in the conventional recording below.

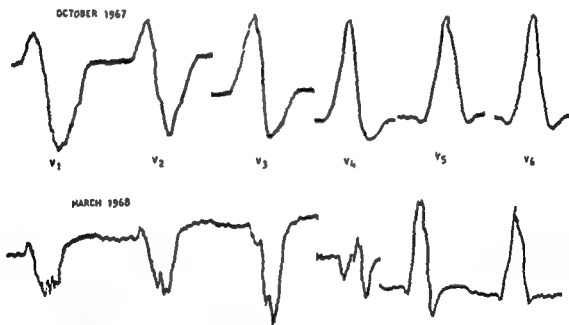


Fig 2 The upper panel consists of high frequency chest leads made in 1967 prior to infarction. The bottom panel shows high frequency records made from the same precordial sites after infarction. After infarction Lead V_4 consists of a QS complex, and V_4 shows obvious low frequency serial change. None of the notching evident in the lower tracing, was present in the conventional electrocardiogram of the same date.

be seen in Table I that 94 per cent of normals have 12 or fewer notches and slurs and 91 per cent of those with heart disease have more than 12 notches and slurs. In Table II 89 per cent of subjects without heart disease have six or less notches and slurs whereas 89 per cent of those with heart disease had more than six notches and slurs. Table III contains the number of notches

(slurs omitted) in the three largest limb leads and in V_1 , V_2 and V_4 . Ninety-two per cent of those without heart disease had three notches or less but 93 per cent of subjects with heart disease had more than three notches. Results from Tables I, II and III were examined statistically. The Median Test² confirmed that each of the three procedures (nine leads, counting

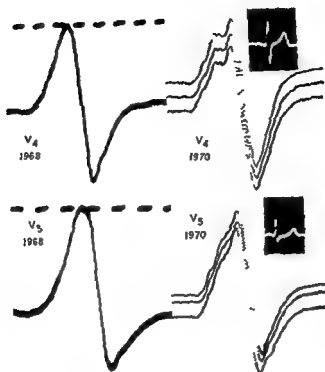


Fig 3 High frequency recording at sites V_4 and V_5 prior to infarction in 1968 and after a well documented episode of myocardial infarction in 1970. The latter records were made by photographing the oscillographic display of a Linc 8 computer. The conventional 12 lead electrocardiogram in 1970 (V_4 and V_5 inverted) was normal.

Table IV Total number of high frequency notches in leads V_4 , V_5 and V_6 together with the total number of notches in the three largest limb leads (Health Evaluation Center)

| Normal | | Abnormal | | | |
|-------------------|--------|-----------------------|--------|--------------|----------------------|
| Number of notches | Normal | Myocardial infarction | Angina | Hypertension | Hypercholesterolemia |
| 0 | 281 | 1 | 2 | 10 | 7 |
| 1 | 268 | 1 | 2 | 8 | 4 |
| 2 | 123 | 2 | 0 | 8 | 5 |
| 3 | 107 | 1 | 1 | 5 | 3 |
| 4 | 50 | 2 | 1 | 2 | 0 |
| 5 | 20 | 1 | 0 | 1 | 0 |
| 6 | 10 | 2 | 1 | 0 | 0 |
| 7 | 6 | 3 | 0 | 0 | 1 |
| 8 | 4 | 1 | 1 | 1 | 0 |
| 9 | 3 | 1 | 0 | 0 | 0 |
| 10 or more | 3 | 9 | 4 | 0 | 0 |
| Totals | 875 | 24 | 12 | 35 | 20 |

Table II Combined number notches and slurs in three largest limb leads and in V_1 , V_2 , and V_3

| Notches and slurs | Normal subjects | Abnormal subjects | Notches and slurs | Normal subjects | Abnormal subjects |
|-------------------|-----------------|-------------------|-------------------|-----------------|-------------------|
| 0 | 6 | 0 | 15 | 0 | 4 |
| 1 | 11 | 0 | 16 | 0 | 4 |
| 2 | 18 | 1 | 17 | 0 | 3 |
| 3 | 17 | 0 | 18 | 0 | 2 |
| 4 | 12 | 2 | 19 | 0 | 7 |
| 5 | 16 | 3 | 20 | 0 | 2 |
| 6 | 9 | 2 | 21 | 0 | 6 |
| 7 | 3 | 4 | 22 | 0 | 1 |
| 8 | 5 | 2 | 23 | 0 | 0 |
| 9 | 1 | 6 | 24 | 0 | 1 |
| 10 | 1 | 4 | 25 | 0 | 1 |
| 11 | 0 | 7 | 26 | 0 | 1 |
| 12 | 0 | 4 | 27 | 0 | 0 |
| 13 | 0 | 5 | 28 | 0 | 0 |
| 14 | 0 | 3 | 29 | 0 | 1 |
| | | | 30 | 0 | 0 |
| | | | Totals | 100 | 16 |

Table III Notches in three largest limb leads and in V_1 , V_2 , and V_3

| Notches | Normal subjects | Abnormal subjects | Notches | Normal subjects | Abnormal subjects |
|---------|-----------------|-------------------|---------|-----------------|-------------------|
| 0 | 38 | 1 | 15 | 0 | 5 |
| 1 | 26 | 2 | 16 | 0 | 2 |
| 2 | 13 | 1 | 17 | 0 | 2 |
| 3 | 15 | 1 | 18 | 0 | 1 |
| 4 | 5 | 3 | 19 | 0 | 2 |
| 5 | 3 | 5 | 20 | 0 | 0 |
| 6 | 0 | 5 | 21 | 0 | 1 |
| 7 | 0 | 7 | 22 | 0 | 0 |
| 8 | 0 | 5 | 23 | 0 | 1 |
| 9 | 0 | 3 | 24 | 0 | 0 |
| 10 | 0 | 3 | 25 | 0 | 0 |
| 11 | 0 | 5 | | | |
| 12 | 0 | 4 | Totals | 100 | 16 |
| 13 | 0 | 10 | | | |
| 14 | 0 | 7 | | | |

cardiograms. Twenty two have continued to be normal. The other two have exhibited serial changes. One subject initially seen in 1966, developed inferior pectoris in 1970. At that time his high frequency electrocardiogram exhibited notching in Leads I, V_R , and V_6 . The second subject entered the study in 1967 at the age of 33 years. In August 1970, he appeared for routine examination. A high frequency electrocardiogram revealed changes not present in 1965 which were most marked in V_4 .

(Fig. 5) Approximately 1 month later he died suddenly of acute myocardial infarction.

Discussion

In 1966 we reported that normal subjects could be separated from subjects with coronary heart disease by counting the total number of notches and slurs in the three largest limb leads and six precordial V leads.⁹ Using criterion of 12 notches and slurs or less to define the normal group we



Fig. 5 The complex at the left is V_4 recorded in 1965. The distorted and notched complex at the right is V_4 obtained in 1970, 6 months later, the patient died of an acute myocardial infarction.

of these hypertensive subjects had left ventricular hypertrophy as noted by conventional electrocardiography with normal heart size judged by routine chest x-ray. None of the 11 had increased notching.

There is evidence that the pathophysiological basis of notching in the electrocardiogram is fragmentation or interruption of the usual smooth path of excitation in the ventricular wall due to patchy necrosis or scarring in the myocardium.^{4,6} Langner and associates⁷ have demonstrated that subpericardial injection of formalin in the intact dog resulted in myocardial lesions and high frequency notching of the QRS complex. Flowers and colleagues¹⁶ have shown that high frequency notching and slurring seen ante mortem in the high frequency electrocardiogram in the orthogonal V and Z leads could be correlated with the site of infarction at post mortem.

If fragmentation of excitation due to necrosis or scarring is indeed one of the mechanisms for the genesis of notching, then our results in instances of two risk factors—hypertension and hypercholesterolemia—can be interpreted to indicate that in cases such as these there may not yet be sufficient myocardial alteration to cause excess notching.

During the present study we encountered seven subjects with angina pectoris who had excess notching in the high frequency electrocardiogram when the conventional electrocardiogram was normal. In an earlier study we reported three cases from our original control group who had excess high frequency notching just prior to development of acute myocardial infarction. Two additional cases of this type have been reported here.

Heart disease is a principal cause of death in this country. A recent study²³ has

indicated that the recognition of coronary heart disease in the preclinical stage at the time of a routine health examination is almost totally inadequate in persons who subsequently died of myocardial infarction. We have shown that high frequency notching in the electrocardiogram is significantly increased in subjects with known coronary heart disease and in some cases appears de novo just before acute myocardial infarction. On the other hand not all patients with coronary artery disease have high frequency notching, as in five of our cases with typical angina pectoris and both a normal conventional electrocardiogram and a normal high frequency electrocardiogram.

Summary

Wide band high frequency electrocardiography employing an expanded time scale and both greater amplitude and a greater frequency response than conventional electrocardiography reveals small notches in the QRS complex which are obscured in the conventional electrocardiogram. A total of more than three high frequency notches in Leads V_1 , V_4 and V_6 plus the notches in the three largest limb leads suggests a high probability of coronary heart disease. This criterion is a modification of one we previously reported in that now fewer leads need be examined and slurs are not counted. In a control group of 100 normal subjects and 76 subjects with coronary artery disease used to develop this new criterion there were 8 false positives (8 per cent) and 5 false negatives (7 percent). The Health Evaluation Center group provided 966 new subjects who were studied using the high frequency technique. 875 were classified as normal. 11 per cent had an abnormal number of notches. Of 24 patients with

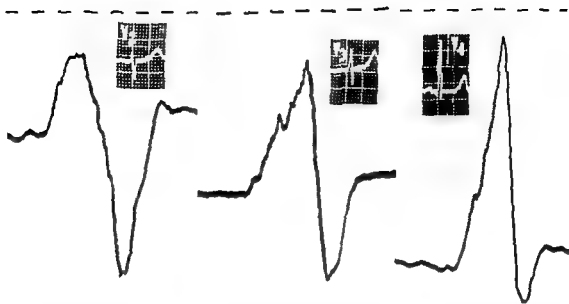


Fig. 4 High frequency electrocardiograms below and conventional electrocardiogram above (Lead V_1 , V_2 and V_4) from a patient with infarct pectoris and 80 per cent occlusion of two coronary arteries on angiography. Notching in the high frequency tracing is self evident. The entire 12 lead conventional electrocardiogram was normal.

observed 6 per cent false positives and 9 per cent false negatives. A restudy of this group of subjects has indicated that adequate separation can be achieved by counting notches alone in V_1 , V_2 and V_4 and in the three largest limb leads. The false positive rate then is 8 per cent while the false negative rate is 7 per cent (Table III). As indicated under *Results*, the confidence level for any of these criteria is high ($P < 0.001$). We feel that the new criterion (counting notches alone in 6 leads) is simpler. The elimination of a count of slurs removes some ambiguity when records are read by visual inspection and also facilitates development of computer programs for interpreting high frequency records. Of interest is the observation that there was no increased notching with age over a ten year period except in three cases who developed myocardial infarction.

Although we have found the counting of notches alone adequate as a screening procedure for reading a single high frequency electrocardiogram recorded from the standard limb and precordial V leads, there may be instances when slurs are significant. Flowers, Horan, and colleagues¹⁷ have found slurs as well as notches significant in their studies which used orthogonal X , Y

and Z leads. In a small study of 20 subjects comparing standard precordial V leads with orthogonal X and Z leads we have found that a given high frequency component appearing as a definite notch in a precordial V lead may appear as a slur in the X or Z lead.²²

The Health Evaluation Center study has enlarged the overall number of subjects in the long term study. This group consists of ambulatory working males who are predominantly normal. The percentage of false positives (11 per cent) is 3 per cent higher than in the group used to develop the present notching criterion. However, follow up for the Health Evaluation Center group was about 30 per cent compared to 100 per cent for the Provident Mutual Life Insurance Company employees who have been followed for a much longer period of time.

Abnormal subjects from the Health Evaluation Center constitute a small group. The percentage of false negatives in the infarct group (24 cases) was 21 per cent. In a group of twelve subjects with a diagnosis of infarct pectoris there were 42 per cent false negatives. In the two high risk groups—hypertension and hypercholesterolemia—no increase in the number of subjects with excess notches or

- 25 Durrer D van Lier A A W and Buller J
 Epicardial and intramural excitation in chronic
 myocardial infarction. *Am HEART J* 68:165
 1964
- 26 Durrer D The human heart some aspects of
 its excitation. *Trans Stud Coll Physicians
 Phila* 33:159 1966
- 27 Langner P H Jr DeMott T and Hussey
 M High fidelity electrocardiography Effects
 of induced localized myocardial injury in the
 dog. *Am HEART J* 71:190 1966
- 28 Schor S S Clark F W Parkhurst I W
 Baker J P and Elsom A V An evaluation
 of the periodic health examination the findings
 in 350 examinees who died. *Ann Intern Med*
 61 999 1964

histories of myocardial infarction, 79 per cent showed excess notching. Of 12 subjects with angina pectoris 7 showed excess notching. Notching in the 35 patients with uncomplicated hypertension and in 20 subjects with hypercholesterolemia was within the normal range. Twenty-four additional Provident Mutual Life Insurance Company employees were added to our long-term study. Of these two developed coronary artery disease. Both exhibited serial changes in the high frequency electrocardiogram.

Wide band electrocardiography using the sum of the notches in the three largest limb leads and the notches in V_1 , V_2 and V_6 is a useful adjunct for the detection of coronary heart disease especially when the conventional electrocardiogram is normal.

We wish to thank Thomas W. Clark MD for his assistance in data collection and Doyle H. Stree BA MA for his help with the statistical analysis. We also wish to acknowledge the technical assistance of Mrs. Freda S. Leiser.

REFERENCES

- Langner P H Jr. The value of high fidelity electrocardiography using the cathode ray oscillograph and an expanded time scale. *Circulation* 5:249 1952.
- Langner P H Jr. High fidelity electrocardiography: further studies including the comparative performance of four different electrocardiographs. *Am Heart J* 45:683 1953.
- Langner P H Jr. Further studies in high fidelity electrocardiography: myocardial infarction. *Circulation* 8:905 1953.
- Langner I H Jr and Geselowitz D B. Characteristics of the frequency spectrum in the normal electrocardiogram and in subjects following myocardial infarction. *Circ Res* 8:577 1960.
- Langner P H Jr, Geselowitz D B and Minsure F T. High frequency components in the electrocardiograms of normal subjects and of patients with coronary heart disease. *Am Heart J* 62:746 1961.
- Langner P H Jr and Geselowitz D B. First derivative of the electrocardiogram. *Circ Res* 10:220 1962.
- Geselowitz D B, Langner P H Jr and Minsure F T. Further studies on the first derivative of the electrocardiogram including instruments available for clinical use. *Am Heart J* 64:805 1962.
- Langner P H Jr. Serial change in high frequency electrocardiograms: a 10 year follow up. Presented at the Cardiac Electrophysiological Group Meeting, Atlantic City, N J, May 31 1964.
- Langner P H Jr and Lauer J A. The relative significance of high frequency and low frequency notching in the electrocardium. *Am Heart J* 71:34 1966.
- Burch G E, Horan I G, Abildskov J A and Cronvich J A. A study of the spatial vectorcardiogram in subjects with posterior myocardial infarction. *Circulation* 12:418 1955.
- Burch G E, Horan I G, Ziskind J and Cronvich J A. Correlative study of post-mortem electrocardiographic and spatial vectorcardiographic data in myocardial infarction. *Circulation* 18:375 1958.
- Trinke F K, Braunstein J R and Zeller D C. Study of high frequency components in electrocardiogram by power spectrum analysis. *Circ Res* 10:340 1962.
- Selvester R H, Rubin H B, Hamlin J A and Pote W W. New quantitative vectorcardiographic criteria for the detection of suspected myocardial infarction in diabetics. *Am Heart J* 75:335 1968.
- Boyle D, Carson P and Hamer J. High frequency electrocardiography in ischemic heart disease. *Br Heart J* 28:539 1966.
- Flowers N C, Horan I G, Thomas J R and Tolleson W J. The importance of high frequency components in the electrocardiogram. *Circulation* 39:531 1969.
- Flower N C, Horan I G, Tolleson W J and Thomas J R. Localization of the site of myocardial scarring in man by high frequency components. *Circulation* 40:977 1969.
- Flowers N C and Horan I G. Diagnostic import of QRS notching in high frequency electrocardiograms of living subject with heart disease. *Circulation* 44:605 1971.
- Reynolds F W, Muller B F, Anderson G J and Muller B F. High frequency components in the electrocardiogram: a comparison study of normals and patients with myocardial disease. *Circulation* 35:195 1967.
- Holcroft J W and Lieberman J. Notching of the QRS complex in high frequency electrocardiograms of normal children and in children with rheumatic fever. *J Electrocardiol* 3:133 1970.
- Minsure F T and Langner P H Jr. A high speed camera for high frequency electrocardiography. *Am Heart J* 67:88 1964.
- Chung S. Small computer analysis of wide band electrocardiograms. Philadelphia December 1960. M.S.E. Thesis University of Pennsylvania.
- Downie N M and Heath R W. Basic statistical methods. 3rd ed. New York 1960. Harper & Row Publishers pp 268 290.
- Langner P H Jr and Minsure F T. High frequency electrocardiography: a comparison of orthogonal vector leads and precordial V leads. Proceedings of the Long Island Jewish Hospital Symposium on Vectorcardiography, Amsterdam 1966. North Holland Publishing Co 147.
- Durrer D, Formigne I, van Duijn R Th, Buller J, van Lier A A W and Meyler R I. The electrocardiogram in normal and some abnormal conditions. In revised human fetal heart and in acute and chronic coronary occlusion. *Am Heart J* 61:303 1961.

struction and the rest had pure valvular stenosis. The age range was from one to 47 years with a mean of 16 years. Patients with left to right shunts at any level were excluded but small to moderate right to left shunts on the atrial level were accepted. Two patients had slight tricuspid regurgitation but none had frank congestive heart failure. Thirty-three of the patients had ECG and VCG recordings made within one week after the hemodynamic investigation. In the other five this interval was about two years but in none of them was there any change in ECG or clinical condition during this period. Three subjects had a QRS duration of 0.11 or 0.12 second in the presence of a conduction defect could not be ruled out. None had QRS duration exceeding 0.12 second. Subjects with the Turner phenotype were excluded.

ECG and VCG. Twelve lead ECGs were recorded with Flema-Schonander-Mingographs of various models all having a high frequency response (linear to 500 Hz) and a high input impedance (exceeding 10 megohms). VCGs were recorded with the axial lead system¹ by means of silver electrodes and a Sanborn 185B amplifier (input impedance about 100 megohms) and a 569B visoscope. The skin was rubbed with acetone before electrode attachment. Ordinary axis orientation was used. Vector loops were photographed in the horizontal, right sagittal and frontal projections with special emphasis on the presentation of the initial dots. The highest frequency response and a beam interruption interval of 25 msec was always used. Scalar X, Y and Z leads were recorded with an Elema-Schonander-Mingograph 34 with the characteristics mentioned at paper speeds of both 50 and 250 mm per second.

X, Y and Z components of each spatial vector were derived through a manual analysis of scalar and planar recordings. The process involved in many instances some degree of judgment as to the identification of vectors in different planes, the main difficulty being that the dots were not identically timed on different pictures. Maxima and minima were derived solely from the scalar tracings. Since the continuous record allowed better identification of optima situated between dots and the

Table 1 Data selected for correlation with right ventricular pressure

I Vectorcardiographic data

- X, Y and Z of each 10 msec QRS vector up to 80 msec
- X, Y and Z of maximal T and ST vector
- QRS maximum and minimum of X, Y and Z
- X minima were separated into early (septal) and secondary (later than 20 msec) minima
- Time from QRS onset to each maximum and minimum QRS duration
- QRS rotation in horizontal plane (counterclockwise = 1 figure of 8 = 2 clockwise = 3)
- Maximal patial QRS vector (MSV)
- Maximal rightward patial vector (MRSV)
- Definition 1 = maximal patial vector pointing to the right of the midline. Definition 2 = spatial magnitude of maximum rightward vector (X min)
- Maximal rightward—anteriorly oriented QRS deflection along the 135 to 315 degree axis in horizontal plane (max RA)
- Maximal leftward—posteriorly oriented QRS deflection parallel with the same axis (max LI)
- Arithmetic sum of RA LI deflections for each 10 msec QRS vector (ΣRA LI)

B Electrocardiographic data

RV₁, SV₁, Ra₁R, R I S I R V₁, SV₁

C Other data

Age

average of several beats could be used.

Hemodynamic data. Right heart catheterization was performed in all patients and right ventricular angiography was made in all except those with a right ventricular pressure below 50 mm Hg. The peak systolic right ventricular pressure was selected as the single hemodynamic variable of interest. This pressure is pathophysiologically and clinically fundamental in this lesion and has previously been shown to be the hemodynamic variable which correlates best with ECG data.^{2,3}

Right ventricular systolic pressure did often vary somewhat during the investigation. The most representative resting pressure was selected by the hemodynamic investigator independently of ECG or VCG data.

Data selection. For each patient 58 items of VCG and ECG data were selected (Table 1). The data were chosen to comprise almost all information present in the QRS complex while ST and T were recorded in less detail. The QRS rotation in the horizontal plane was coded as figure-of 8 when even

Prediction of right ventricular systolic pressure in pulmonary stenosis from combined vectorcardiographic data

Knut Rasmussen MD*
Sigmund J Sørland MD
Oslo, Norway

Although heart catheterization and angiocardiography have become increasingly precise and safe procedures, it still is and will be an important goal in cardiology to extract as much information as possible from noninvasive techniques. The best such method for the diagnosis of ventricular hypertrophy has been electrocardiography (ECG) in one form or another. Isolated right ventricular pressure overload as seen in pulmonary stenosis with intact ventricular septum is well suited for the testing of further progress in this field. The lesion is hemodynamically simple and the easily obtainable right ventricular peak systolic pressure is commonly accepted as the best index of degree of disease.¹

The hypothesis underlying most work done with the ECG prediction of hemodynamic data has been that there is a fundamental relationship between the functional load, the degree of anatomical changes induced by this load and some properties of the electrical forces created within these anatomical structures.¹ One has therefore sought for the components of these forces which give the best prediction of pressure or flow. Generally three meth-

ods are available in the efforts to improve the results obtained with conventional ECG. (1) An orthogonal vectorcardiographic (VCG) system may be substituted for the LCC.^{2,3} (2) The dipole model may be changed to a more complex physical model involving multiple dipoles⁴ or multipoles.⁵ Although the first of these methods recently has been shown to yield promising results, these methods will for a long time remain investigational procedures. (3) The information obtained through LCC or VCG could be used to better advantage through the combination of several measurements.

This report presents the results of our efforts along lines (1) and (3). The multilead system has been used for VCG measurements and a multiple regression analysis has been made in order to select the best data combination for pressure prediction.

Materials and methods

Patients Thirty-eight unselected patients with a diagnosis of unoperated pulmonary stenosis with intact ventricular septum were included in the study. Four patients had a pure infundibular stenosis, two had combined valvular and infundibular ob-

From Medical Department B and the Diagnostic Department, University Hospital, Rikshospitalet, Oslo, Norway.
Received for publication Oct. 30, 1972.

Reprint requests to Dr. Knut Rasmussen, Rikshospitalet, Med. Dept. B, Oslo, Norway.

*Research fellow. The Norwegian Research Council for Scientific Health Insurance.

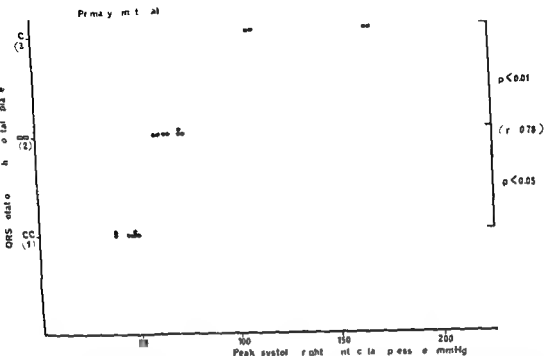


Fig 1 Relationship between direction of QRS inscription in horizontal plane and right ventricular pressure

new equilibrium has had time to be established.^{11,12} In three cases the time between the pressure measurement and the VCG recording was more than half a year with some clinical or ECC signs of progression. The first five patients had small left to right shunts on the ventricular level (below 2% per cent of pulmonary flow) and one of them had in addition an intermittent third degree A-V block. Thus this material was less homogeneous than the primary one but was nevertheless thought to be useful for the evaluation of the derived formula.

Results

Altogether 40 of the 58 independent variables proved to be significantly correlated with right ventricular systolic pressure at the 5 per cent level and 33 of the correlations were significant also on the 1 per cent level. The best of these individual correlation coefficients are presented in Table II. As seen the best individual predictor was found to be the classical VCG variable of direction of QRS rotation in the horizontal plane. The calculation of a correlation coefficient for this variable is how-

ever not quite correct since it is not continuously distributed. The individual data for QRS rotation pressure relationships are shown in Fig 1.

Another closely related and continuously distributed index namely (max RA-max LP) had the same correlation coefficient (0.78) (Fig 2). We have found this measurement to be of some value because the variation around the regression line is smallest in the lower pressure range where the practical importance of an index of severity is greatest. For a sample of this size the 95 per cent confidence interval of this correlation coefficient is from 0.61 to 0.88.

It should further be noted from Table II that the best conventional FCC variable came out as No. 6 and that the second definition of VRSV was somewhat better than the first and more commonly applied definition. The new indices based on RA-LP relationships were not particularly useful compared with more commonly used variables. Time variables did generally yield poorer results than the voltage variables.

The multiple stepwise regression calcu-

Table II Correlation* coefficient between individual ICG and VCG measurements and right ventricular systolic pressure

| A | Variable | r |
|----|-------------------------------|-------|
| 1 | QRS rotation horizontal plane | 0.78 |
| 2 | Max RA | 0.74 |
| 3 | $\Gamma /$ | 0.74 |
| 4 | $S I /$ | 0.74 |
| 5 | MRSV definition 2 | 0.73 |
| 6 | $R V_1$ | 0.72 |
| 7 | $S I$ | 0.72 |
| 8 | $/ \text{min}$ | 0.71 |
| 9 | $/ 50 \text{ msec}$ | -0.71 |
| 10 | $\Sigma R A I I$ | 0.68 |
| 11 | MRSV definition 1 | 0.68 |
| 12 | Σmin_2 | 0.68 |
| 13 | $\Delta S V$ | 0.67 |
| 14 | $/ 60 \text{ msec}$ | -0.67 |
| 15 | Time $/ \text{min}$ | 0.66 |
| 16 | $/ 40 \text{ msec}$ | -0.64 |
| 17 | ΓV_1 | -0.63 |
| 18 | $\Delta 60 \text{ msec}$ | -0.61 |
| 19 | $\Delta 70 \text{ msec}$ | -0.57 |
| 20 | $R A V_R$ | 0.56 |
| 21 | $/ 70 \text{ msec}$ | -0.55 |
| 22 | $Z \text{ max}$ | -0.52 |
| 23 | $S V_1$ | -0.51 |
| 24 | Max LP | -0.51 |
| 25 | $\Delta 70 \text{ msec}$ | 0.50 |
| 26 | $\Delta 60 \text{ msec}$ | 0.50 |
| 27 | $\Sigma T V_1$ | -0.48 |
| 28 | $\Delta 80 \text{ msec}$ | 0.47 |
| 29 | $\Delta 80 \text{ msec}$ | 0.46 |
| 30 | $\Delta 50 \text{ msec}$ | -0.45 |
| 31 | $Z 80 \text{ msec}$ | -0.45 |
| 32 | $\Delta 50 \text{ msec}$ | 0.42 |
| 33 | $S V_1$ | 0.42 |

*All correlations are significant ($p < 0.01$ or less). The variables are ranked according to the height of the coefficient.

the slightest part of the loop had rotation opposite to the main loop. Both of the two possible definitions of the much used maximal rightward spatial vector (MRSV) were included.

With increasing right ventricular hypertrophy the QRS loop shifts gradually from a posterior leftward to an anterior rightward position in average approximately around the 315 to 135 degree axis in the horizontal plane. An effort was therefore made to quantitate this shift through the measuring of the maximal anterior rightward (AR) and posterior leftward (PL) deflections parallel with this axis. The shift

of the total QRS loop was quantitated through the derivation of the sum of these deflections for each 10 msec vector.

Although the study was not primarily designed to compare LCC and VCC, some of the most used LCC measurements for the diagnosis of right ventricular hypertrophy were also selected. Age was used as the single non-ECC variable.

Data treatment. The 58 items of data on each patient were entered as independent variables and the peak systolic right ventricular pressure as the dependent variable in a multiple regression computer program (BIVID 09)⁸ using a CDC 3600 computer. This program is essentially a combined step up and step down procedure¹⁰ starting with the best individual predictor and at each step including the new predictor which gives the greatest reduction in the rest variance in terms of having the largest F value. Predictors which lose their significance during the procedure are discarded. F values of 5.0 were chosen as limits for both entering and removal of variables corresponding to a minimum significance level of about 5 per cent. Through this process the formula of the order

$$\text{Pressure} = a + b_1x_1 + b_2x_2 + \dots + b_nx_n$$

which gives the best possible prediction of pressure in this material is computed.

In addition to the Cartesian coordinates a computer run was also made with derived spherical coordinates (spatial magnitude, azimuth and elevation). Because of the slight skewness in the distribution of right ventricular pressure in the material logarithmic transformations of the variables were also tested.

Secondary material. Any regression equation derived from a limited sample may give poorer results in a secondary sample. Therefore a secondary material consisting of 19 unselected patients with pulmonary stenosis as their main lesion was sampled in order to test the results. Five patients satisfied the inclusion criteria of the primary material. Six patients had been operated on for their lesion more than one year previously and a new hemodynamic investigation had been made. Previous studies indicate that ECC pressure correlations are unchanged in such patients if a

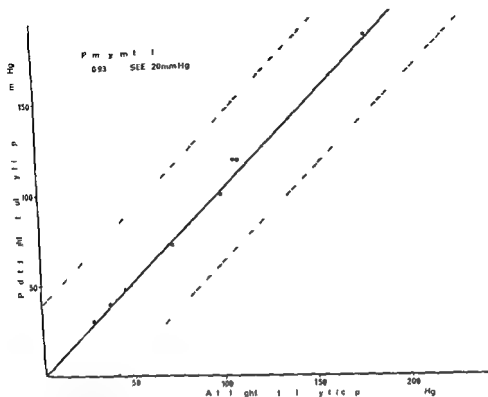


Fig. 3 Relationship between actual right ventricular systolic pressure and the pressure computed by means of the equation described in the text

selected except for the exclusion of subjects with the Turner phenotype. This material is the first of its type in which this group has been explicitly excluded. This would be expected to result in improved correlations and has definitely done so in this material. Patients with Turner phenotype and normal chromosomes constitute about seven per cent of the population with pulmonary stenosis.⁷

The age range in this material is large compared to many previously reported; the mean age is also comparatively high.

ECG VCG technique. Both the recording technique and the manual derivation of instantaneous vectors contain possible errors. The quality of the data could probably have been somewhat improved through the use of computer wave recognition and data sampling.¹² The greater error in the measurement of instantaneous vectors than of maxima and minima may possibly be partly responsible for the comparatively low diagnostic performance of the former. This work may therefore be regarded as representative of what can be

accomplished through manual ECG and VCG measurements which in most centers will remain the only method for many years.

Data selection. An almost infinite number of VCG variables and combinations may be derived and analyzed. The possibility that other indices could improve the results is always there, but it seems unlikely that the improvement would be great.

Because the number of ECG variables tested was so small the results have limited value in the comparison of the optimal diagnostic performance of ECG vs VCG. It is possible that individual or combined ECG variables could be found in this material that might give as good results as the VCG variables.

It is also possible that the results could be improved by taking other hemodynamic parameters into consideration, for instance left ventricular systolic pressure.

Data treatment. The choice of the coefficient of correlation and the standard error of estimate as the sole criterion of diagnostic performance in this and pre-

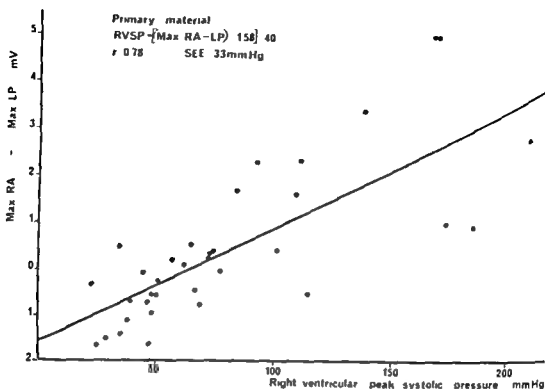


Fig 2 Relationship between max RA-max LP index and right ventricular systolic pressure. The scatter around the regression line is least in the 50 to 80 mm Hg region.

lation proceeded through five steps at each step a new variable was introduced into the equation. The following formula was derived:

$$\text{RVSP} = -57.6 + 15.4 \text{ ST Z} + 21.3 \Delta \text{ max} + 62.4 \Delta \text{ min} + 35 \text{ QRS rot} + 1.4 \text{ age}$$

Through the use of this formula the correlation coefficient between actual and predicted pressure increased from 0.78 to 0.93; the 95 per cent confidence interval of the latter being from 0.87 to 0.96. The standard error of the pressure estimate (SEE) was reduced from 33 to 20 mm Hg (Fig 3).

All the coefficients in the equation had standard errors of less than one third of their absolute values.

The use of spherical coordinates in the regression analysis did not improve the results. Again horizontal plane QRS rotation was selected as the first step; the next being the spatial magnitude of S1 and maximal T vector. The individual vectors' spatial magnitude correlated much better than the angles with pressure. Logarithmic transformations gave much poorer results.

An effort to combine the seven ECG variables and age into a formula was unsuccessful, since it produced only two significant steps.

As would be expected from the less rigid

patient selection, most of the individual predictors showed poorer results when applied to the secondary material (QRS rotation $r = 0.69$, RV1 $r = 0.68$). Others showed better results than in the primary material (MRSV $r = 0.91$, max RA $r = 0.89$, S1 $r = 0.79$). The derived equation gave a somewhat—but not much—poorer prediction than in the primary material with a correlation coefficient of 0.88 (95 per cent confidence interval 0.71 to 0.91) (Fig 4). Two patients in the secondary material fell outside the ± 2 SEE of the primary material: one above and one below the regression line. The one above had a ventricular septal defect with a variable left to right shunt which may have added significantly to the right ventricular load; and the one below had A-V block during the catheterization (heart rate 44 per minute) and the pressure measured may therefore have been higher than usual. Thus the deviations may to some extent be explained by the different sample selection.

Discussion

Patient selection. This is an important aspect in this kind of work, since the exclusion of a few atypical patients may falsely improve the results considerably. The present material is in this regard un-

1 me 80
1 mb r 3

the square root of the number of patients. This requirement is satisfied in the present work. With increased sample size the multiple regression method may be applied with greater benefit and confidence. Even then the value of the formula derived can be properly tested only on future samples as attempted here.

Results Individual correlation coefficients
With the above mentioned sources of error in mind, some of the results will be compared with previous ones.

In this study single VCC variables were found to have a small superiority over single ECG variables. This is in contrast to Combar and co-workers² who found a much larger benefit of Frank VCG and also in some contrast to Borun and Sipin³ (cube and Frank VCC) who found no benefit at all. A moderate diagnostic improvement resulting from the substitution of VCG for ECG was also found by Witham and associates⁴ (Helm system) and Hoft and associates⁵ (Frank system). No conclusion can be drawn regarding the superiority of one VCC system or another in this type of diagnosis from these studies. The present work is however the first of this kind in which the axial lead system has been used and the results are well comparable with those obtained with other orthogonal systems.

The degree of correlation found here for several simple ECG and VCG indices (RV₁, S₁, V₁RSV) is in good accordance with several previous reports^{2,4,17,6}. The best individual correlations have been published by Witham and associates⁴ but they were using rather uncommon VCG variables and the material was small.

The high diagnostic performance of the QRS rotation in the horizontal plane that was found is at some variance with previous results with other lead systems. For the cube system this is probably caused by too high sensitivity to slight increments in right ventricular pressure.^{19,21} Clockwise rotation of the loop seems to occur at about the same pressure level in the Frank and the axial lead systems² but the transition from counterclockwise to clockwise rotation is possibly more abrupt with the Frank system.² This is however partly dependent on the exact definition of loop rotation applied.

Voltage maximum and minimum were found to be definitely more firmly related to right ventricular pressure than in the time intervals used to reach these optima (Table II). This is contrary to the proposition recently made by Arntzenius² to use such time intervals in the diagnosis of ventricular hypertrophy. In addition his method may be difficult to apply because the deflection points may have no precise definition in abnormal loop rotations. Both the time interval method and the use of single vector magnitudes in the diagnosis of hypertrophy are based on the partly invalid hypothesis that single instantaneous vectors may be generated in only one part of the heart.

Recent work on the VCC diagnosis of right ventricular hypertrophy has been dominated by the use of the VRSV as an index. The results from the primary material (Table II) indicate that this dominant role is not altogether justified. On the other hand VRSV was the best predictor in the smaller secondary material. The new index based on anterior rightward loop dislocation did not prove to be particularly useful.

In recent VCC studies either Cartesian or spherical coordinates have been used; this divergence has been one of the many in this field which has decreased the applicability of the results. This study indicates that no information is lost by using the less sophisticated Cartesian coordinates.

Benefit from multiple regression analysis
In the present study a statistically significant and potentially practically important improvement of pressure VCC correlations was provided through the combination of several VCG data. The derived equation is simple and rapid to apply; in addition to scalar VCG leads one needs only a photograph of the horizontal plane QRS loop.

Two groups of workers have previously to some degree applied the principle of predictor combination for estimating right ventricular pressure in pulmonary stenosis. Witham and associates⁴ selected eight promising variables for a multiple regression analysis, thereby restricting the factors which could be taken into account. They obtained a formula which included seven primary variables; a large number considering the small sample of 21 patients. When comparing with the best individual variable only a small improvement was found.

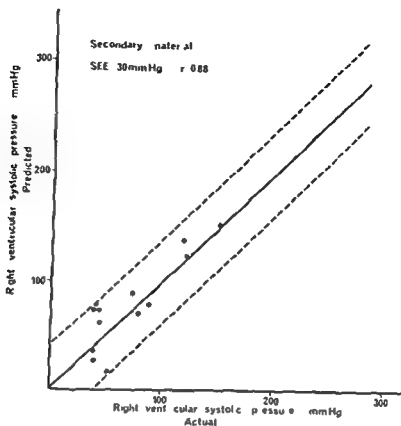


Fig. 4 Relation between actual and equation predicted systolic right ventricular pressure in the secondary material

vious works^{2,3} may be disputed. This choice implies that all pressure regions are treated as equally important while the practical importance of the correlations is greater in the 50 to 100 mm Hg region. The derived equation is actually somewhat better in the upper pressure region.

Linear regression analysis does in a strict sense require a normal distribution of data. This is not completely satisfied in this material, but better than in several previous ones.^{2,4}

The most important limitation in this and most of the previous studies^{2,4} is the small sample size. This causes the confidence intervals of the coefficients to become so large that reliable conclusions concerning the relative diagnostic value of different variables and combinations can often not be drawn. This factor has often not been properly considered and may explain most if not all of the differences in results for instance regarding the superiority of ECG or VCG.¹⁴ This applies also to most of the differences found in this study. The comparison of correlation coefficients and standard errors of estimate from different materials is still more misleading.

As demonstrated by the large number of significant individual predictors the situation dealt with is statistically characterized by a large number of measurements which are highly correlated with right ventricular pressure and which also have a high degree of intercorrelation. The problem in the multiple regression analysis is to select the variable which at each step gives most additional knowledge. Beyond the first step this is usually not the variable with the best individual performance. The variable that is selected is also dependent on very small differences in F values. A new computation from a slightly different sample may lead to the selection of an entirely different set of variables but probably providing about the same amount of information as the first set. Thus too much weight should not be placed upon the selection of variables that have appeared in the final equation.

Uncritical application of results from multiple regression analysis of a sample to the population may lead to serious errors.¹⁵ Gumbor, Klingeman and Pipberger¹⁶ advocate that the number of variables included in a multivariate formula should not exceed

catheterized patients. This situation is somewhat different since previous recordings from the same patient may serve as control data and thereby increase the diagnostic accuracy. Changes from time to time in VCC pressure relationship may be assumed to occur parallel with the regression line. (3) The effect of surgery may similarly be evaluated thus contributing to the decision of whether a postoperative catheterization should be made.

In patients with right ventricular hypertrophy caused by various other kinds of congenital and acquired heart disease the pressure VCG relationship will usually be different particularly when a flow load is added to a pressure load. However if the VCG is interpreted not in terms of pressure but of anatomy the results may possibly have value also in other conditions.

Summary

In 38 patients with isolated unoperated pulmonary stenosis a systematic search was made for optimal VCG criteria for the prediction of peak systolic right ventricular pressure. Fifty VCC measurements, seven ECC measurements and age of each patient were entered into a stepwise multiple regression computer program.

The best individual predictors were found to be the QRS loop rotation in the horizontal plane and the closely related QRS dislocation along the 135 to 315 degree horizontal plane axis ($r = 0.78$). Five VCG criteria were better than the best ECG criterion ($R V_1$, $r = 0.72$). Thirty three of the 58 variables showed significant correlations with the pressure ($p < 0.01$). Since the confidence intervals are large with this sample size and degree of correlation conclusions regarding the superiority of one predictor vs another should be drawn with great care.

The multivariate equation selected by the computer involved four VCG variables and age; this improved the correlation coefficient to 0.93. This improvement from data combination is larger than in previous studies, probably because all variables were given equal opportunity to enter the equation.

The results were tested on a secondary sample of 19 patients with pulmonary stenosis as their main cardiac lesion. Although

this sample was less homogeneous the formula derived pressure estimates remained reasonably good ($r = 0.88$). The study suggests that the diagnostic power of FCC and VCG could be increased through the proper combination of easily obtainable measurements.

REFERENCES

1. Hugenoltz P G and Cambo A R. Effect of chronically increased ventricular pressure on electrical forces of the heart. *Circulation* 40:511 1969.
2. Cambo A R, Hugenoltz P G and Nadas A S. Corrected (Frank) uncorrected (Cube) and standard electrocardiographic lead systems in recording augmented right ventricular forces in right ventricular hypertension. *Br Heart J* 28:62 1966.
3. Witham A C, Rainey R L and Edmonds J H Jr. Prediction of right ventricular pressure in pulmonic stenosis from sponge vector cardiogram and electrocardiogram. *Am Heart J* 75:187 1968.
4. Holt J H, Barnard A C L, Lynn M S and Kramer J O. A study of the human heart as a multiple dipole electrical source. III. Diagnosis and quantitation of right ventricular hypertrophy. *Circulation* 40:711 1969.
5. Borun E R and Sapin S O. Comparison of Frank Cube and conventional electrocardiographic data with hemodynamic measurements in children with pulmonic stenosis. In: *Vector cardiography 2* (Proceedings of the 1st International Symposium on Vectorcardiography, Amsterdam 1971, North Holland Publishing Company) p. 309.
6. Arthur H M, Geselowitz D B, Briller S A and Trost R F. Quadrupole components of the human surface electrocardiogram. *Am Heart J* 83:663 1972.
7. Rasmussen K and Serland S. Electrocardiogram and vectorcardiogram in Turner phenotype with normal chromosomes and pulmonary stenosis. *Br Heart J* In press.
8. McFee R and Parungao A. An orthogonal lead system for clinical electrocardiography. *Am Heart J* 62:93 1961.
9. Elroyson M A. Multiple regression analysis in Ralston and Wilf editors. *Mathematical methods for digital computers*. New York 1960. John Wiley & Sons Inc. Part V 17.
10. Armitage P. *Statistical methods in medical research*. Oxford 1971. Blackwell Scientific Publications. Chap. 10 p. 314.
11. Kahn M, Bleifer S B, Grishman A and Donoso E. The vectorcardiogram and electrocardiogram before and after valvulotomy for pulmonic stenosis. *Am Heart J* 58:327 1959.
12. Hanson J S, Ikkes D, Crafoord C and Oventen C O. Results of surgery for congenital pulmonary stenosis. *Circulation* 18:588 1958.
13. Caceres C A and Hochberg H M. Per

with reduction of standard error of pressure estimate from 19 to 15 mm Hg

Borun and Sipin⁶ studied a much larger sample of 90 patients with Frank and cube VCG and convention of ECG. They found the best single variable to be $R V_1$ ($r = 0.79$, $SEE = 19$ mm Hg). Adding one or two variables (S_x or $R_x/[R_x + S_x]$) improved the correlation coefficient to 0.83 and 0.84. They apparently did not use a computer program to select the optimal data combination.

Thus the main difference between these two reports and the present is that in the latter all 58 variables have been given an equal opportunity of being included in the equation. This may probably explain most of the larger benefit from data combination.

Secondary material. As discussed any correlation, selected because it is good, will tend to decrease when a second sample is analyzed. Multiple regression equations are particularly vulnerable in this respect. The presently supplied secondary material was not ideally suited because of the heterogeneity, but it was the only sample available. The great changes in correlation for several of the individual predictors illustrate the sample dependency of these results. The persisting good correlation between actual and formula predicted pressure, with most of the data falling within ± 2 SEE of the regression line indicate that the formula really reflects properties of the pulmonary stenosis population as such and that it may be of practical help in future work with VCG in this group. Although two individual variables yielded results about as good as the formula in the secondary material, the formula is definitely better when both materials are considered and showed a better sample stability.

General discussion. Even if one assumes a complete correspondence between physiology, anatomy, and ECG, the correlations between these entities must be restricted by the repeat variabilities of the measurements involved. The repeat variability of ECG and VCG data is largely known²² but data on the reliability of right ventricular pressure measurements are as far as we know not available. However, even during the same investigation the pressure may vary by several millimeters of mercury. Therefore, it seems doubtful if SEE of such

correlations can be reduced to below 10 mm Hg.

Another source of persisting variation around the regression line is the factor of obstructive infundibular pulmonary stenosis. In this condition the resting pressure may be low but it may increase steeply at small levels of exercise. ECG and VCG may correctly show changes compatible with advanced right ventricular hypertrophy and are therefore a better indicator of degree of disease than an isolated pressure measurement.

Different left ventricular pressures with consequent differences in left ventricular electrical forces will probably also decrease the correlations; this factor may partly have been responsible for bringing the age into the equation.

The main conclusion which may be drawn from the present work is that there is a substantial benefit to be gained from combining VCG data in the diagnosis of right ventricular hypertrophy. After the formula has been developed, all that is required is conventional VCG measurements and a short time for calculation. This time is well spent in this group of patients where important decisions regarding prognosis, catheterization, and operation are to be made. It is possible that the results achieved in this way may be as dependable as those obtained by the use of complex physical models involving a large number of measurements and complicated computer calculations.⁴ Before spending too much energy on these interesting lines of research one should ascertain that more easily obtainable data are used optimally.

The practical importance of the results lies in the improved ability to estimate the degree of pulmonary stenosis in cases where this diagnosis is fairly certain. The three typical situations of use are as follows: (1) In the previously uncatheterized patient the regression equations may be helpful in establishing if heart catheterization is indicated and if operation should be planned at an early stage. The practical value of the pressure estimate in this situation is restricted by the fact that catheterization is also made in order to confirm the diagnosis and to exclude the presence of additional defects. (2) The method may be used to evaluate progression in previously

catheterized patients. This situation is somewhat different since previous recordings from the same patient may serve as control data and thereby increase the diagnostic accuracy. Changes from time to time in VCG pressure relationship may be assumed to occur parallel with the regression line. (3) The effect of surgery may similarly be evaluated thus contributing to the decision of whether a postoperative catheterization should be made.

In patients with right ventricular hypertrophy caused by various other kinds of congenital and acquired heart disease the pressure VCG relationship will usually be different particularly when a flow load is added to a pressure load. However if the VCG is interpreted not in terms of pressure but of anatomy the results may possibly have value also in other conditions.

Summary

In 38 patients with isolated unoperated pulmonary stenosis a systematic search was made for optimal VCG criteria for the prediction of peak systolic right ventricular pressure. Fifty VCG measurements, seven ECG measurements and age of each patient were entered into a stepwise multiple regression computer program.

The best individual predictors were found to be the QRS loop rotation in the horizontal plane and the closely related QRS dislocation along the 135 to 315 degree horizontal plane axis ($r = 0.78$). Five VCG criteria were better than the best ECG criterion (R_{V1} , $r = 0.72$). Thirty three of the 38 variables showed significant correlations with the pressure ($p < 0.01$). Since the confidence intervals are large with this sample size and degree of correlation conclusions regarding the superiority of one predictor vs another should be drawn with great care.

The multivariate equation selected by the computer involved four VCG variables and age. This improved the correlation coefficient to 0.93. This improvement from data combination is larger than in previous studies probably because all variables were given equal opportunity to enter the equation.

The results were tested on a secondary sample of 19 patients with pulmonary stenosis as their main cardiac lesion. Although

this sample was less homogeneous the formula-derived pressure estimates remained reasonably good ($r = 0.88$). The study suggests that the diagnostic power of ECG and VCG could be increased through the proper combination of easily obtainable measurements.

REFERENCES

1. Hugenholz J G and Gamboa R. Effect of chronically increased ventricular pressure on electrical forces of the heart. *Circulation* 30: 311, 1964.
2. Gamboa R, Hugenholz J G and Nadas A S. Corrected (Frank) uncorrected (Cube) and standard electrocardiographic lead systems in recording augmented right ventricular forces in right ventricular hypertension. *Br Heart J* 28: 62, 1966.
3. Witham A C, Rainey R L and Edmonds J H Jr. Prediction of right ventricular pressure in pulmonary stenosis from sponge vector cardiogram and electrocardiogram. *Am Heart J* 72: 187, 1968.
4. Holt J H, Barnard A C L, Lynn M S and Kramer J O. A study of the human heart as a multiple dipole electrical source. III. Diagnosis and quantitation of right ventricular hypertrophy. *Circulation* 40: 711, 1969.
5. Boron E R and Sapin S O. Comparison of Frank Cube and conventional electrocardiographic data with hemodynamic measurements in children with pulmonary stenosis. In: *Vector cardiography 2. Proceedings of the 1st International Symposium on Vectorcardiography, Amsterdam 1971*. North Holland Publishing Company, p 560.
6. Arthur R M, Geselowitz D B, Briller S A and Tost R F. Quadrupole components of the human surface electrocardiogram. *Am Heart J* 63: 663, 1972.
7. Rasmussen H and Sorland S. Electrocardiogram and vectorcardiogram in Turner phenotype with normal chromosomes and pulmonary stenosis. *Br Heart J* In press.
8. McFee R and Parungao A. An orthogonal lead system for clinical electrocardiography. *Am Heart J* 62: 93, 1961.
9. Ekstrom M A. Multiple regression analysis. In: *Kalton and Wolf editors. Mathematical methods for digital computers*. New York 1960. John Wiley & Sons, Inc. Part V: 17, p 191.
10. Armitage P. Statistical methods in medical research. Oxford 1971. Blackwell Scientific Publications. Chap 10, p 314.
11. Kahn M, Blefer S H, Grishman A and Donoso E. The vectorcardiogram and electrocardiogram before and after valvulotomy for pulmonary stenosis. *Am Heart J* 58: 327, 1959.
12. Hanson J S, Hnos D, Crawford C and Owenfora C O. Results of surgery for congenital pulmonary stenosis. *Circulation* 111: 588, 1958.
13. Caceres C A and Hochberg H M. Per

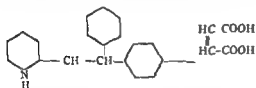
- formance of the computer and physician in the analysis of the electrocardiogram. *Am Heart J* 79:439, 1970
14. Pipberger H V, Schneidermann M A and Klingeman J D. The love-it first night effect in research. *Circulation* 38:872, 1968
 15. Cooley W W and Lohnes P R. Multivariate procedures for the behavioral sciences. New York, 1962. John Wiley & Sons Inc. p. 35
 16. Gimboa R, Klingeman J D and Pipberger H V. Computer diagnosis of biventricular hypertrophy from the orthogonal vectorcardiogram. *Circulation* 39:72, 1969
 17. Basingthwaite J B, Parkin T W, DuShane J W, Wood I H and Burchell H B. The electrocardiographic and hemodynamic findings in pulmonary stenosis with intact ventricular septum. *Circulation* 28:893, 1963
 18. Scherlis I, Koenker R J and Lee Y C. Pulmonary stenosis. Electrocardiographic vectorcardiographic and catheterization data. *Circulation* 28:288, 1963
 19. Yabum J H, Dufano M J and Toor M. Pulmonic stenosis. A clinical assessment of severity. *Am J Cardiol* 5:44, 1960
 20. Ellison R C and Restierux N J. Quantitation of ventricular hypertrophy and hemodynamic load with vectorcardiogram. *Br Heart J* 35:599, 1973
 21. String R H, Hugenholz P G, Liebman J and Nadas A. The vectorcardiogram in pulmonary stenosis. Correlation with the hemodynamic state in patients with and without ventricular septal defect. *Am J Cardiol* 12:758, 1963
 22. Arntzenius A C. A physiological approach to vectorcardiographic recognition of ventricular hypertrophy. *Cardiovasc Res* 1:105, 1970
 23. Willemis J L, Poblete I F and Pipberger H V. Day-to-day variation of the normal orthogonal electrocardiogram and vectorcardiogram. *Circulation* 45:1057, 1972

The action of perhexiline maleate in patients with angina

C M Morgans M1 MB MRCP
J Russell Rees MA MD MRCP
Bristol England

Perhexiline maleate is a new antianginal drug found to be effective in recent double blind trials in America and the United Kingdom. In a majority of the patients there was a substantial reduction in anginal attacks and in glyceryl trinitrate consumption.^{1,2}

Perhexiline maleate closely resembles a predecessor hexadiline which was reported on by Rowe and associates.³ Its chemical name is 2-(2,2-dicyclohexylethyl) piperidine maleate; it has the following structural formula:



In animals the drug induces systemic arterial vasodilatation, increased coronary blood flow, reduced heart rate and increased pulmonary compliance.^{4,5}

We report here a double blind clinical study of perhexiline undertaken to assess its effect on heart rate and blood pressure at rest, on graded exercise and at the point of angina. Respiratory function and daily walking were also measured.

Patients and methods

Eleven men with typical angina pectoris of frequent occurrence were invited to the study. Their ages ranged from 45 to 67 with a mean of 53. Length of history ranged from six months to 12 years with a mean of five years. Five had normal resting electrocardiograms (ECG); two showed horizontal ST segment depression, one an old anterior and one an old posterior myocardial infarct. One showed complete left and the last complete right bundle branch block. None of the ECGs altered during the trial. There was no recent change in the pattern of angina pectoris in any patient and no recent myocardial infarction. At the time of study none showed left ventricular failure.

The study consisted of a two week control period during which all tablets except glyceryl trinitrate were discontinued, followed by a four week trial period. The trial was double blind; the patient taking either perhexiline maleate (200 mg) at 9 AM and 3 PM or placebo tablets identical in appearance and taste each for two weeks by random selection during the trial period. Each patient attended for exercise tests on a bicycle ergometer on three occasions at fortnightly intervals starting at the end of the control period.

Received for publication June 19, 1973

Revised for publication December 19, 1973

Revised for publication December 19, 1973

Deposited with the British Library Document Supply Centre, Boston, MA 01905, U.S.A.

1186 No 3 pp 379-383 Sept 1973

American Heart Journal 329

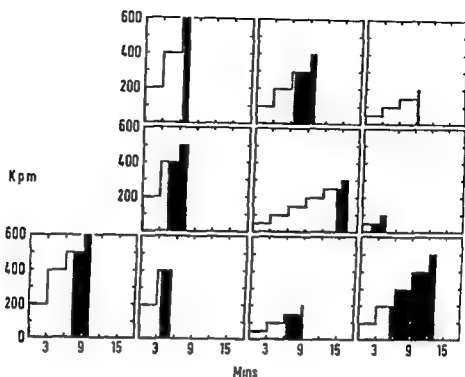


Fig 1 Performance on drug vs placebo. Hatched sections represent exercise tolerance on placebo. Dark sections represent increase in exercise tolerance on perhexiline. End point in each case is angina.

On the test days the patient was instructed not to smoke and not to drink tea or coffee after 10 a.m. He had a light lunch at 12.30 p.m. and attended the Cardiac Department in the afternoon. Patients were asked to avoid taking glyceryl trinitrate on test days if the drug was taken, the tests were abandoned.

On arrival the patient rested in an examination room for ten minutes. He was then examined, the pulse rate over one minute was measured and also the blood pressure by sphygmomanometry. The patient walked through to the study room where an ECG was recorded. Remaining supine he was attached to a bicycle ergometer (Elema Schonander type AM 368). Lead V_3 was monitored by oscilloscope to which a recorder was attached. Five minutes later pulse rate and blood pressure were again recorded.

The patient then cycled at 60 revolutions per minute in the supine position, the load increasing every three minutes. At each load blood pressure was recorded after two minutes and the ECG after two and a half minutes. The heart rate at each load was measured as the mean of 20 beats. At the first visit the loading was adjusted until angina developed in less than 15 minutes.

At each subsequent visit the patient cycled to the point of angina three times and stopped five seconds later, the end point heart rate being measured during this interval. Three loading regimes were used: 50 k.p.m. increasing by 50 k.p.m. increments; 100 k.p.m. increasing by 100 k.p.m. increments; and 200 k.p.m. increasing to 400, 500 and 600 k.p.m.

Room temperature was kept between 73° and 78° F. It did not vary by more than 3° F for any patient. Forced vital capacity and forced expiratory volume were recorded on a Vitalograph. The patient was instructed in the use of a pedometer and the number of miles walked during the trial was recorded.

Results

One of the 11 patients discontinued his tablets subsequently found to be perhexiline after four days of the second period of the trial because of dizziness. The results analyzed are those from the other ten patients.

When the code was broken after the study and all measurements had been completed it was found that six patients had taken perhexiline first and four placebo. Exercise tolerance was greater in

Table I Work performance on placebo and perhexiline

| Patient | Placebo (k p m × time) | Perhexiline (k p m × time) | Percentage increase on perhexiline |
|---------|------------------------------|----------------------------------|--|
| 1 | 2 030 | 2 630 | 30 |
| 2 | 1 120 | 2 200 | 96 |
| 3 | 910 | 1 026 | 13 |
| 4 | 1 147 | 2 183 | 90 |
| 5 | 2 033 | 2 563 | 26 |
| 6 | 99 | 272 | 175 |
| 7 | 2 500 | 4 190 | 68 |
| 8 | 947 | 1 620 | 71 |
| 9 | 472 | 913 | 100 |
| 10 | 807 | 3 733 | 363 |
| Mean | 1 206 | 2 136 | 77 |

all cases on perhexiline than on placebo ($p < 0.01$). The increase in the total amount of work performed (product of work load and time) varied from 13 to 362 per cent with a mean of 77 per cent (Fig. 1 Table I). Two patients who had perhexiline before placebo had an exercise tolerance on placebo less than after the control period.

In the first test of each study day (Table II) resting heart rates on perhexiline were significantly lower than on placebo but on placebo they were higher in this test than in subsequent ones probably due to anxiety. In the second and third tests this difference diminished as resting heart rates on placebo approached those on perhexiline which remained unchanged.

Heart rates at the highest work load achieved on placebo were compared with those at the same work load on perhexiline. Eight of the ten showed lower rates with perhexiline and the difference in the whole group is significant for each test (Table III).

Heart rates at the point of angina were not significantly different and did not vary by more than ten beats per minute except in the two patients who did not have lower exercising heart rates on perhexiline (Patients 1 and 8 Table IV). Their heart rates at the point of angina were 11 and 30 beats per minute higher on perhexiline.

Blood pressure readings used in the analysis were the mean of those recorded in the study room during the second and third tests. No significant difference be-

Table II Resting heart rates (mean values for 10 patients)

| Test | Placebo | Perhexiline | Differences between perhexiline and placebo |
|------|---------|-------------|---|
| I | 77.9 | 69.0 | -8.9 $p < 0.01$ |
| II | 73.2 | 69.3 | -3.9 $p = NS$ |
| III | 72.3 | 69.0 | -3.3 $p = NS$ |

Table III Heart rates per minute at the highest comparable work load (mean values for 10 patients)

| Test | Control | Placebo | Perhexiline | Difference p value |
|-----------------|---------|---------|---------------------|-----------------------|
| I Load varying | 104.0 | 96.8 | -7.2 $p < 0.05$ | |
| II Load varying | 106.0 | 97.0 | -9.0 $p < 0.001$ | |
| III | 103.5 | 103.5 | 97.5 $p < 0.05$ | |

Table IV Heart rates per minute at the point of angina

| Patient | Perhexiline | Placebo |
|---------|-------------|---------|
| 1 | 139 | 128 |
| 2 | 111 | 105 |
| 3 | 106 | 110 |
| 4 | 109 | 103 |
| 5 | 111 | 112 |
| 6 | 81 | 87 |
| 7 | 95 | 104 |
| 8 | 150 | 170 |
| 9 | 103 | 110 |
| 10 | 110 | 116 |
| Mean | 111 | 109 |

tween resting or exercising systolic or diastolic blood pressures was found between the values recorded on perhexiline and those on placebo (Table V). The product of systolic blood pressure and heart rate at the highest comparable work load was signifi-

Table V Blood pressure results (mean values for 10 patients)

| | Placebo | Perhexiline | Difference | Significance |
|--|---------|-------------|------------|-------------------|
| Resting systolic (mm Hg) | 136.8 | 136.3 | -0.5 | $p = \text{N.S.}$ |
| Systolic at highest comparable load (mm Hg) | 157.9 | 153.0 | -4.9 | $p = \text{N.S.}$ |
| Resting diastolic (mm Hg) | 83.8 | 85.9 | +2.1 | $p = \text{N.S.}$ |
| Diastolic at highest comparable load (mm Hg) | 97.7 | 93.7 | -4.0 | $p = \text{N.S.}$ |
| Systolic at highest load (mm Hg) \times heart rate | 16,798 | 14,985 | -1,813 | $p < 0.01$ |

cantly lower on perhexiline ($p < 0.01$)

The forced expiratory volume in one second as a percentage of the forced vital capacity was greater than 75 per cent on placebo and perhexiline in all but one patient in whom it was 69 per cent on placebo and 75 per cent on perhexiline. Pedometry revealed no evidence that the patients walked further when taking perhexiline rather than placebo, even though all showed improvement in exercise tolerance.

Discussion

Double blind studies may be imperilled by side effects or the success or the drug under study. It is difficult to avoid this problem and in our trial it was possible to anticipate prior to analysis that the patient who withdrew was on the drug at the time, and in each of the others the correct order of administration for drug and placebo.

The action of the drug was evident in all but the trial was designed to study its effects on heart rate and blood pressure and it was found that blood pressure was unaltered by perhexiline but that heart rates were changed.

Resting rates were 9 per minute slower than on placebo in the first of the three tests, but this statistically significant difference all but disappeared in the next two tests as resting pulse rates on placebo fell. It is possible that the patients were more relaxed in later tests on the same afternoon and that perhexiline abolished the resting tachycardia. If mild anxiety or aware that they were probably on perhexiline they were more confident from the start.

Perhexiline also slowed the exercising

heart rate in all tests and at maximum comparable work load prior to developing angina an average over all rate reduction of 7 per minute was observed.

At the point of angina however, heart rates were the same with perhexiline and placebo supporting the simple explanation that the drug delays angina by a reduction in the rate of rise in heart rate resulting from exercise and possibly emotion.

There are no comparable reports on the effect of the drug on heart rate at the time of angina. Three other studies on heart rate at rest and on exercise have however, been made. Grupp and associates⁹ gave 100 mg q.d.s. of perhexiline to healthy volunteers tested on a treadmill; they found no change in resting heart rates but a reduction in the tachycardia produced by exercise without any effect on blood pressure. Winsor² gave the drug in the same dosage to angina patients also tested on a treadmill. He similarly found no reduction in resting heart rate but a significant fall in exercise heart rate averaging 22 per minute more than that found by us. Lyon and associates² found no consistent change in either resting or postexercise heart rate.

Perhexiline has a direct effect on atrial muscle and is not thought to act through beta blockade.¹⁰ There are differences between the effects we found and those reported from the use of beta blocking drugs in patients with angina. Using propranolol, Prichard and Gillam¹¹ found a dose related reduction in resting and exercising heart rates. Similar reports with alprenolol came from Sealey and associates¹² and Sowton and Smithen.¹³ At effective dosage the reduction was of greater degree than found by us with perhexiline but more important

all three report a reduction in heart rate at the point of angina in contrast to our findings

There are some points to be made against the premise that perhexiline acts solely by delaying the rise in exercise heart rate (1) This effect is modest (2) Perhexiline is sometimes effective in patients not helped by beta blockers known to cause greater reductions in heart rate (3) Perhexiline helped two of our patients without any effect on heart rates at rest or on exercise enabling greater exercise tolerance and heart rates 11 and 30 per minute faster than on placebo at the point of angina If this is not due to a natural remission we must look for another effect of perhexiline

Although powerful coronary vasodilator drugs such as dipyridamole have proved ineffective in angina glyceryl trinitrate has been shown to increase coronary flow in experimental myocardial infarction¹⁴ It remains a possibility that the modest coronary vasodilator action of perhexiline may contribute to its action It has yet to be shown how perhexiline compares with the beta blocking drugs and whether their effects are additive

Summary

In patients with angina a double blind trial of the action of perhexiline maleate at rest and during exercise on a bicycle ergometer has shown a significant increase in exercise tolerance and a reduction in exercising heart rate but no reduction in heart rate at rest or at the point of angina and no change in blood pressure

REFERENCES

- 1 Burns Cox C J Chandrasekhar K P Ikram
H Pe rce T H Picher J Quinlan C D M

- and Rees J R Clinical evaluation of perhexiline maleate in patients with angina pectoris *Br Med J* 4:586 1971
- 2 Hershleifer I Perhexiline maleate in the treatment of angina pectoris *Curr Ther Res* 11:99 1969
- 3 Lyon L J Nevins M A Fisch S and Henry S Perhexiline maleate in treatment of angina pectoris *Lancet* 1:1272 1971
- 4 Martins de Oliveira J and de Almeida Amado N J Perhexiline maleate in the treatment of angina pectoris *Hospital* 77:1511 1970
- 5 Winsor T Clinical evaluation of perhexiline maleate *Clin Pharmacol Ther* 11:85 1970
- 6 Rowe G G Alonso S Boake W C Castillo C A Lugo J E and Crumpton C W Systemic and coronary hemodynamic effects of hexadylamine *Proc Soc Exp Biol Med* 112:545 1961
- 7 Hudak W J Lewis R E and Kuhn W L Cardiovascular pharmacology of perhexiline *J Pharmacol Exp Ther* 131:1371 1970
- 8 Cho Y W Beley M and Wiado D M Pharmacology of a new antianginal drug Perhexiline I Coronary circulation and myocardial metabolism *Chest* 58:377 1970
- 9 Grupp I L Bunde C A and Grupp G Effects of perhexiline maleate on exercise-induced tachycardia *J Clin Pharmacol* 10:317 1970
- 10 Matsuo S Cho Y W and Wiado D M Pharmacology of a new antianginal drug Perhexiline II Heart rate and transmembrane potential of cardiac tissue *Chest* 58:581 1970
- 11 Frichard B N S and Gillam I M S Assessment of propranolol in angina pectoris *Br Heart J* 33:473 1971
- 12 Sealey H J Liljedal J Byberg G and Abbad B Acute effects of oral alprenolol on exercise tolerance in patients with angina pectoris *Br Heart J* 33:481 1971
- 13 Sowton E and Smithen C Double blind three-dose trial of oral alprenolol in angina pectoris *Br Heart J* 33:601 1971
- 14 Pedding V J and Rees J R Early changes in collateral flow following coronary artery ligation The role of the sympathetic nervous system *Cardiovasc Res* 2:219 1968

Table V Blood pressure results (mean values for 10 patients)

| | Placebo | Perhexiline | Difference | Significance |
|--|---------|-------------|------------|--------------|
| Resting systolic (mm Hg) | 136.8 | 136.3 | -0.5 | p = NS |
| Systolic at highest comparable load (mm Hg) | 157.9 | 153.0 | -4.9 | p = NS |
| Resting diastolic (mm Hg) | 83.8 | 85.9 | +2.1 | p = NS |
| Diastolic at highest comparable load (mm Hg) | 97.7 | 93.7 | -4.0 | p = NS |
| Systolic at highest load (mm Hg) X heart rate | 16.798 | 14.985 | -1.813 | p = < 0.01 |

cantly lower on perhexiline ($p < 0.01$)

The forced expiratory volume in one second is a percentage of the forced vital capacity was greater than 75 per cent on placebo and perhexiline in all but one patient, in whom it was 69 per cent on placebo and 75 per cent on perhexiline. Pedometry revealed no evidence that the patients walked farther when taking perhexiline rather than placebo, even though all showed improvement in exercise tolerance.

Discussion

Double blind studies may be imperilled by side effects or the success of the drug under study. It is difficult to avoid this problem, and in our trial it was possible to anticipate prior to analysis that the patient who withdrew was on the drug at the time and in each of the others the correct order of administration for drug and placebo.

The action of the drug was evident in all but the trial was designed to study its effects on heart rate and blood pressure, and it was found that blood pressure was unaltered by perhexiline but that heart rates were changed.

Resting rates were 9 per minute slower than on placebo in the first of the three tests, but this statistically significant difference all but disappeared in the next two tests as resting pulse rates on placebo fell. It is possible that the patients were more relaxed in later tests on the same afternoon and that perhexiline abolished the resting tachycardia of mild anxiety or aware that they were probably on perhexiline, they were more confident from the start. Perhexiline also slowed the exercising

heart rate in all tests and at maximum comparable work load prior to developing angina, an average over all rate reduction of 7 per minute was observed.

At the point of angina, however, heart rates were the same with perhexiline and placebo supporting the simple explanation that the drug delays angina by a reduction in the rate of rise in heart rate resulting from exercise and possibly emotion.

There are no comparable reports on the effect of the drug on heart rate at the time of angina. Three other studies on heart rate at rest and on exercise have, however, been made. Grupp and associates⁹ gave 100 mg q.d.s. of perhexiline to healthy volunteers tested on a treadmill; they found no change in resting heart rates but a reduction in the tachycardia produced by exercise, without any effect on blood pressure. Winsor⁸ gave the drug in the same dosage to angina patients also tested on a treadmill. He similarly found no reduction in resting heart rate but a significant fall in exercise heart rate averaging 22 per minute more than that found by us. Lyon and associates³ found no consistent change in either resting or postexercise heart rate.

Perhexiline has a direct effect on atrial muscle and is not thought to act through beta blockade.¹⁰ There are differences between the effects we found and those reported from the use of beta blocking drugs in patients with angina. Using propranolol, Prichard and Callam¹¹ found a dose related reduction in resting and exercising heart rates. Similar reports with alprenolol came from Senley and associates¹² and Sowton and Smith¹³. At effective dosage the reduction was of greater degree than found by us with perhexiline but, more important

all three report a reduction in heart rate at the point of angina in contrast to our findings

There are some points to be made against the premise that perhexiline acts solely by delaying the rise in exercise heart rate (1) This effect is modest (2) Perhexiline is sometimes effective in patients not helped by beta blockers known to cause greater reductions in heart rate (3) Perhexiline helped two of our patients without any effect on heart rates at rest or on exercise enabling greater exercise tolerance and heart rates 11 and 30 per minute faster than on placebo at the point of angina If this is not due to a natural remission we must look for another effect of perhexiline

Although powerful coronary vasodilator drugs such as dipyridamole have proved ineffective in angina glyceryl trinitrate has been shown to increase coronary flow in experimental myocardial infarction¹¹ It remains a possibility that the modest coronary vasodilator action of perhexiline may contribute to its action It has yet to be shown how perhexiline compares with the beta blocking drugs and whether their effects are additive

Summary

In patients with angina a double blind trial of the action of perhexiline maleate at rest and during exercise on a bicycle ergometer has shown a significant increase in exercise tolerance and a reduction in exercising heart rate but no reduction in heart rate at rest or at the point of angina and no change in blood pressure

REFERENCES

- 1 Burns-Cox C J Chandrasekhar K P Ikram H Tierce T H Pilcher J Quinlan C D M

- and Rees J R Clinical evaluation of perhexiline maleate in patients with angina pectoris *Br Med J* 4 586 1971
- 2 Hirschleifer I Perhexiline maleate in the treatment of angina pectoris *Curr Ther Res* 11 99 1969
- 3 Lyon I J Nevins M A Fisch S and Henry S Perhexiline maleate in treatment of angina pectoris *Lancet* 1 1272 1971
- 4 Martins de Oliveira J and de Almeida Amado N J Perhexiline maleate in the treatment of angina pectoris *Hospital* 7 1511 1970
- 5 Winsor T Clinical evaluation of perhexiline maleate *Clin Pharmacol Ther* 11 85 1970
- 6 Rowe G G Alfonso S Boake W C Castillo C A Lugo J E and Crumpton C W Systemic and coronary hemodynamic effects of hexadylamine *Proc Soc Exp Biol Med* 112 545 1963
- 7 Hodak W J Lewis R E and Kuhn W L Cardiovascular pharmacology of perhexiline *J Pharmacol Exp Ther* 137 371 1970
- 8 Cho Y W Beley M and Aviado D M Pharmacology of a new antianginal drug Perhexiline I Coronary circulation and myocardial metabolism *Chest* 58:577 1970
- 9 Grupp I L Bunde C A and Grupp G Effects of perhexiline maleate on exercise-induced tachycardia *J Clin Pharmacol* 10 312 1970
- 10 Matsuo S Cho Y W and Aviado D M Pharmacology of a new antianginal drug Perhexiline II Heart rate and transmembrane potential of cardiac tissue *Chest* 58 581 1970
- 11 Prichard D N S and Gillam J M S Assessment of propranolol in angina pectoris *Br Heart J* 33 473 1971
- 12 Sealey H J Liljedal J Byberg G and Abbad B Acute effects of oral alprenolol on exercise tolerance in patients with angina pectoris *Br Heart J* 33 601 1971
- 13 Sowton E and Smithen C Double blind three dose trial of oral alprenolol in angina pectoris *Br Heart J* 33 601 1971
- 14 Redding V J and Rees J R Early changes in collateral flow following coronary artery ligation The role of the sympathetic nervous system *Cardiovasc Res* 2 219 1968

Tuberculoma of the heart

Report of 9 cases

O P Kapoor, MD MRCP (Ed)

E Mascarenhas, MD*

M M Rananavare, MD

R K Gadgil MD, PhD, FRCP (London)

Bombay, India

A tubercular granuloma (tuberculoma) in the myocardium is decidedly rare. Autopsy studies carried out prior to introduction of specific antituberculous therapy, state that the myocardium is involved in less than 0.30 per cent of patients dying from tuberculosis.^{1,2} The introduction of specific chemotherapy has diminished the incidence even further³ and over the last decade, these lesions have become the subject of single case reports.⁴⁻⁶ Myocardial tuberculomas have been directly responsible for death in a few instances. The cause of death in these patients has been conduction disturbances,³ ventricular rupture,⁴ coronary artery occlusion,⁵ obstruction to pulmonary blood flow,⁶ and impaired myocardial contractility.⁹ Advances in surgical techniques have now made these lesions potentially correctable,⁶ and heart block secondary to a critical lesion in the conduction system³ may benefit from electrical pacing. Optimal preoperative planning requires an awareness of the disease and precise delineation of the lesion by cardiac catheterization and angiographic techniques.⁵

Since tuberculomas have seldom been

diagnosed during life, the antemortem clinical features of nine patients with myocardial tuberculomas collected from the autopsy records of J J Hospital Bombay, India, are presented. The literature pertaining to morbidity and mortality rate of tuberculomas located in various regions of the heart is reviewed.

Material and methods

During a period of ten years (1961 to 1970) 9333 autopsies were performed at the J J Group of Hospitals, Bombay, India, and nine cases of myocardial tuberculomas were found. The incidence of tuberculosis in the autopsy material was 7.9 per cent. Table I summarizes some of the pathologic features observed in these 9 patients. The lesions were located in all regions of the heart and varied in size from pea to lemon size. The number of tuberculomas in a single heart varied from 1 to 7. The nodules were white to yellowish in color, had a firm consistency, presented an irregular surface, and often had a caseous center. Microscopically these nodules were generally composed of a central area of caseation. At the periphery numerous

From the Department of Medicine and Pathology, Grant Medical College, J J Hospital, Bombay, India.
Received for publication Nov 10, 1972.

Reprint requests to O P Kapoor MD, Khadi Bhavan, Jeevan Udhyog, 278 D N Road, Bombay 1, India.

*Present address: Division of Cardiology, Mount Sinai Hospital, Miami Beach, Fla 33140.



Fig 1 Microphotograph of a tuberculoma showing caseous material in the center surrounded by Langhans giant cells epithelioid cells and lymphocytes (Hematoxylin and eosin. Original magnification $\times 480$)

Table 1 Table of nine patients with myocardial tuberculomas

| Serial no | Patient initials | Age/sex | Site of myocardial tuberculoma | Endocarditis | Myocarditis | Pericarditis | Probable primary focus |
|-----------|------------------|---------|--------------------------------|--------------|-------------|--------------|---|
| 1 | S P | 8/F | Right atrium | + | | | Hilar glands lungs |
| 2 | S D | 52/M | Right atrium | | | + | Hilar glands lungs |
| 3 | M A | 3/M | Right atrium | | | | Hilar glands lungs |
| 4 | R R | 42/M | Right atrium | | | + | Hilar glands lungs |
| 5 | P R | 37/M | Right atrium | | | | Hilar glands lungs |
| 6 | C R | 55/M | Left atrium | | + | | Hilar glands lungs |
| 7 | M M | 5/M | Left ventricle | | | | Miliary tuberculosis (?) blood borne |
| 8 | R C | 0/1 | Multiple | | | + | Hilar glands lungs |
| 9 | F D | 45/M | Multiple | | | + | Hilar glands lungs |

Acid fast bacilli de tubed

follicles consisting of epithelioid cells giant cells and lymphocytes were seen. In some instances the outer zone consisted of an area of epithelioid and large round cells (Fig 1). In all patients an attempt was made to identify acid fast bacilli by application of Ziel-Neelsen stain; however tubercle bacilli were identified in only 2 instances. In the remaining patients the lesions were presumed to be tubercular because of the characteristic histologic picture along with an active focus of tuberculosis in the body. A pericarditis of varying degree in either the visceral or parietal component of the pericardial sac was observed in four instances. The endocardium remained intact in all but two patients. A

diffuse interstitial and parenchymal myocarditis was noted in one patient. Histologically this myocardium revealed an increase in fibrous tissue cellular infiltration and a moderate increase in giant cells.

A resume of the pertinent clinical and pathologic features in the nine patients with myocardial tuberculomas follows. A comment is made on the signs and symptoms that may be produced by tuberculomas located in various regions of the heart.

Tuberculomas of the right atrium

Case 1 An 8-year old girl was admitted for investigation of fever and puffiness of the face of a month's

Tuberculoma of the heart

Report of 9 cases

O P Kapoor, MD MRCP (Ed)

E Mascarenhas MD*

M M Ranana-ware, MD

R K Gadgil MD PhD FRCP (London)

Bombay India

A tubercular granuloma (tuberculoma) in the myocardium is decidedly rare. Autopsy studies carried out prior to introduction of specific antituberculous therapy state that the myocardium is involved in less than 0.30 per cent of patients dying from tuberculosis.^{1,2} The introduction of specific chemotherapy has diminished the incidence even further,³ and over the last decade these lesions have become the subject of single case reports.⁴⁻⁸ Myocardial tuberculomas have been directly responsible for death in a few instances. The cause of death in these patients has been conduction disturbances,⁴ ventricular rupture,⁶ coronary artery occlusion,⁶ obstruction to pulmonary blood flow,⁸ and impaired myocardial contractility.⁹ Advances in surgical techniques have now made these lesions potentially correctable,⁵ and heart block secondary to a critical lesion in the conduction system⁹ may benefit from electrical pacing. Optimal preoperative planning requires an awareness of the disease and precise delineation of the lesion by cardiac catheterization and angiographic techniques.⁵

Since tuberculomas have seldom been

diagnosed during life the antemortem clinical features of nine patients with myocardial tuberculomas collected from the autopsy records of J J Hospital, Bombay India are presented. The literature pertaining to morbidity and mortality rate of tuberculomas located in various regions of the heart is reviewed.

Material and methods

During a period of ten years (1961 to 1970) 9333 autopsies were performed at the J J Group of Hospitals Bombay India, and nine cases of myocardial tuberculomas were found. The incidence of tuberculosis in the autopsy material was 7.9 per cent. Table I summarizes some of the pathologic features observed in these 9 patients. The lesions were located in all regions of the heart and varied in size from pea to lemon size. The number of tuberculomas in a single heart varied from 1 to 7. The nodules were white to yellowish in color and a firm consistency presented an irregular surface and often had a caseous center. Microscopically these nodules were generally composed of a central area of caseation. At the periphery numerous

From the Department of Medicine and Pathology Grant Medical College J J Hospital Bombay India
Received for publication Nov 10 1972

Reprint requests to O P Kapoor MD Khadi Blavan Jeevan Udhog 278 H N Road Bombay 1 India
*Present address Division of Cardiology Mount Sinai Hospital Miami Beach Fla 33140



Fig 3 Case 2 Large caseous tumor in the right atrium



Fig 4 A through C Cases 3, 4, and 5 Tuberculomas in the right ventricular outflow tract (A and B) and anterior right ventricular myocardium (C)

tubercular complex and bilateral hilar tuberculous

Case 4 A 42 year old man was admitted to the Tuberculosis Ward for therapy of pulmonary tuberculosis. On cardiac auscultation a Grade 3/6 ejection systolic murmur was heard at the left sternal border. The second heart sound was normally split with A_2 greater than P_2 . The remainder of the cardiovascular examination was normal. There was no cardiac failure. The patient died a few days later from tuberculosis. At autopsy a large tuberculoma measuring 4 cm, in length 2 cm in width and 1 cm in depth was seen in the anterior wall of the right ventricle. A cut-section revealed a cavity composed of necrotic material with a thick fibrotic wall (Fig

4B). Acid fast bacilli were recovered from this lesion.

Case 5 The heart of a 37 year old man who died on arrival at the hospital showed a tuberculoma 2 cm by 2 cm by 1 cm in the anterior wall of the right ventricle (Fig 4C). The rest of the heart was normal. An active tubercular lesion was present in the right upper lung.

Comment In this series (5 patients, two of whom had multiple tuberculoma) tuberculomas were most commonly seen in the right ventricle. This observation is consistent with the findings of Weigert¹¹ and Ziegler¹² who considered the right ventricular myocardium to be the most common site, especially in the region of the right conus arteriosus (Weigert tubercle). In two of our patients large tuberculomas



Fig 2 Case 1 Open right atrium showing an ulcerating tuberculous mass with thrombus formation at the SVC RA junction

duration. Physical examination revealed facial edema, bilateral nonpulsatile neck vein engorgement and clubbing of the nailbeds. The lymph glands in the left axilla were enlarged and matted. Signs of a cavitating pulmonary lesion were detected in the right lung. The cardiac examination was essentially normal except for the presence of a continuous murmur over the right second and third interspace. A chest x ray confirmed the presence of a cavitating lesion in the right upper lung field. The cardiac silhouette was normal. A diagnosis of pulmonary tuberculosis with mediastinal compression was made following identification of acid fast bacilli in the sputum. Anti tuberculous therapy was instituted; unfortunately the patient expired after five days of a massive bout of hemoptysis.

The heart at autopsy showed a spherical mass 1 cm in diameter in the right upper atrium that projected into the lumen of the superior vena cava. A thrombus had developed over this mass and obstructed the flow of blood from the superior vena cava into the right atrium (Fig 2). The tuberculous nature of this lesion was established by the characteristic granulation tissue and presence of acid fast bacilli on microscopic examination.

Case 2 The heart of a 52 year old man dying of bilateral pulmonary tuberculosis showed a large mass which filled the entire right atrial appendage (Fig 3). The cardiac lesion was not suspected prior to death and retrospectively the only clue to the lesion may have been the presence of long standing atrial fibrillation.

Comment Anderson¹⁰ in his review of 72 patients with myocardial tuberculomas found the right atrium to be the most frequent site of involvement. This was felt to be so because of the frequent involvement of the right mediastinal lymph nodes with spread to the right atrial myocardium. Horn and Saphir¹ remarked about the rarity of endocardial

ulceration. In Case 1 the endocardial continuity was disrupted with subsequent thrombus formation which extended into the SVC giving rise to a superior vena cava obstruction syndrome. A similar case has not been previously reported. In Case 2 the tuberculous mass occupied a significant cross section of the right atrial cavity. Such gross lesions may seriously interfere with atrial contraction as cited by Adamson.⁸ Both patients with right atrial tuberculomas had atrial fibrillation. Gouley and associates¹¹ have commented that the occurrence of atrial ectopic beats in a patient with tuberculosis should alert one to the possibility of involvement of the myocardium. The etiology of atrial arrhythmias may be secondary to irritability produced by the inflammatory process in the atrium or to destruction of the SA node¹² by tuberculous pathology. Even though detailed histologic studies of the SA node were not carried out in Case 1 the location of the tuberculous mass at the SVC RA junction makes it interesting to speculate that the sinus node was perhaps infiltrated by tuberculous granulation tissue.

Tuberculomas of the right ventricle

Case 3 A three year old boy was admitted in a debilitated state with a history of fever of fifteen days. On examination the child was anemic and malnourished. The lung fields were clear to percussion and auscultation. Examination of the heart revealed a Grade 3/6 ejection systolic murmur over the pulmonary area and along the left sternal border. The second heart sound was normally split with diminished intensity of the pulmonary component. A diagnosis of pulmonary valve stenosis was made on the basis of the cardiac auscultatory findings and the etiology of fever was under investigation. The child died 48 hours following admission. At necropsy a lemon sized mass was noted in the right ventricular outflow tract (Fig 4-1). The lungs showed a primary

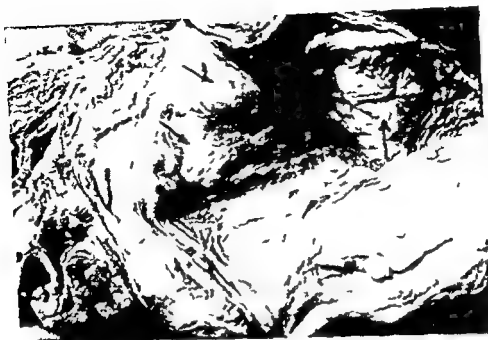


Fig 6 Case 8 Open left ventricle showing multiple tuberculomas in the interventricular septum.



Fig 7 Case 9 Two tuberculous nodules in the wall of the right ventricle and left ventricular apex

appendage anterior right ventricular wall right ventricular infundibular area and apex of the left ventricle. The interventricular septum housed three tuberculomas (Fig 6) the diameter of the largest one being 2.1 cm. There was an associated tubercular

pericarditis and an active tubercular focus in the left upper lung.

Case 9 The heart of a 45 year old man who died of chronic heart failure showed an area of localized pericarditis around the left ventricular apex. A large tuberculoma (3 cm by 2 cm by 1 cm) was present in the anterior wall of the right ventricle and a second lesion (1 cm by 1 cm by 0.5 cm) at the apex of the left ventricle (Fig 7). The lungs showed chronic congestive changes and active tuberculosis of the right upper lung.

Comment Extensive involvement of the intra-ventricular septum was associated with complete A-V block in a case reported by Kinare and Deshmukh.⁸ A detailed study of the conduction system in this case showed marked destruction of the A-V node and His bundle by the tubercular process. In Case 8 besides the tubercular cardiac lesions there was no other demonstrable etiology for the patient's sudden and unexpected demise. Even though detailed studies of the conduction system were not carried out in this instance, the sudden death in this patient may have been as a result of a lethal disturbance of cardiac conduction or rhythm.

Multiple tuberculomas were seen in two of our patients. According to Anders¹³ it is not uncommon to see multiple tuberculomas in a single heart two to five being a fair average. The greatest number recorded is 18 in a case of Fontoyot.¹⁴

Summary

Nine patients with tuberculomas of the heart selected from autopsy material over a ten year period are presented. The signs and symptoms produced by tuberculomas located in various regions of the heart are



Fig 5 Case 6 Circumscribed tuberculous nodule in the wall of left atrium

were situated in the right ventricular outflow tract and had the auscultatory findings of right ventricular outflow tract obstruction. Rawls and colleagues⁴ reported a case similar to our patients in which a large myocardial abscess in a 36-year old woman with tuberculous lymphadenopathy, partially obstructed the right ventricular outflow tract. The clinically suspected hemodynamic alterations were confirmed by cardiac catheterization and right ventricular angiography. The patient had an excellent response to isoniazid and Para amino-salicylic acid and to surgical drainage of the abscess.

The presence of myocardial tuberculomas in patients with tuberculosis may also be suggested by the presence of right bundle branch block pattern or right heart strain on the electrocardiogram.⁵ Even though none of the lesions in this series had perforated, tuberculomas have been known to perforate⁶ into a cardiac chamber or into the pericardial cavity, thus producing widespread dissemination, acute tamponade or sudden death.

Tuberculoma of the left atrium and myocarditis

Case 6 A 55 year old man was seen in the Emergency Room with acute pulmonary edema. He looked severely anemic and had a thready pulse with a regular tachycardia of 150 beats per minute. A chronic suppurating ulcer was noticed over the left mandibular area. The patient died before complete examination could be carried out. At necropsy both lung fields showed fibrocaceous tuberculosis most marked in the left lower lobe. The heart was diffusely dilated and hypertrophied. A peanut sized

nodule was seen projecting from the wall of the left atrium into the cavity (Fig 5). The myocardium showed extensive scarring, lymphocytic infiltration and giant cells. Numerous tubercular granulomas were observed in the left atrial mass, the mandibular ulcer and the lung.

Comment Small tuberculomas in the left atrium may be detected at autopsy with no prior clinical evidence referable to the heart.^{11,12} However, larger lesions in this location have seriously obstructed blood flow through the left atrium. The cases of Maurocordatus¹³ and Townsend¹⁴ showed nodules of the left atrium so large that blood flow in the pulmonary veins was obstructed with resulting hemorrhage in the lungs and death from a phlebotomy.

The etiology of acute pulmonary edema in Case 6 was probably secondary to diffuse myocarditis. Horn and Saphir¹⁵ have cautioned against the diagnosis of tuberculous myocarditis in hearts with excessive connective tissue with a moderate amount of giant cells in the absence of cavitation or acid fast bacilli. Since similar changes will be accounted for by a previous diphtheria, rheumatic fever, scarlet fever, coronary artery disease, etc. The myocarditis in our patient is therefore presumed to be tubercular in etiology because of the presence of acid fast bacilli in other areas of the body.

Tuberculomas of the left ventricle

Case 7 Examination of the heart of a 5 year old boy who died of an automobile accident showed an elongated grayish yellow tumor like mass 3.5 cm by 1.3 cm by 1.2 cm extending from the left ventricular apex and deforming the posterior leaflet of the mitral valve. The mass added 1.2 cm to the thickness of the left ventricular wall and was sub-endocardial throughout its course. Microscopically the mass was composed of numerous granulomas with central caseation. Miliary tubercles were widely distributed in the lungs and liver.

Comment Although Ravasi¹⁶ regarded the left ventricle as the main site of myocardial tuberculous involvement, it has seldom been found as the sole cardiac lesion by others.^{10,11} The reason for this is perhaps because the usual manner of spread of tuberculosis to the heart is from the right mediastinal lymph nodes.¹⁰ In Case 7 the mediastinal lymph nodes were not involved and the patient had generalized miliary tuberculosis. It is conceivable that in this particular instance the myocardial lesion was secondary to a blood borne infection. However, it should also be borne in mind that myocardial tuberculomas may be the means of disseminating this disease and setting up a generalized miliary invasion as observed in one instance by Gouley and colleagues.¹¹ The intact endocardium in our case suggests that the myocardial lesion was the result rather than the cause of generalized miliary tuberculosis.

Multiple tuberculomas and tuberculomas of the intraventricular septum

Case 8 The heart of a 40-year old man who died suddenly and unexpectedly showed seven large tuberculomas. The e were located in the right atrial

Total anomalous pulmonary venous drainage with ventricular septal defect

Carl N. Steeg MD

Kent Ellis MD

Weldon M. Gersony MD

New York, N.Y.

The association of a large ventricular septal defect with total anomalous pulmonary venous drainage has been reported only rarely.^{1,2} The clinical implications of this combination of lesions have not been previously emphasized. Three patients with total anomalous pulmonary venous drainage and ventricular septal defect have been documented at this institution within a one year period. One infant had additional pulmonic stenosis. All presented with clinical findings compatible with large left to right shunts. The diagnosis was made at cardiac catheterization and angiocardiology in two patients but in the third infant the ventricular septal defect was not discovered until repeat catheterization following repair of the total anomalous pulmonary venous drainage.

Incomplete diagnosis may result in serious errors in management. The intent of this report is to outline an approach to diagnosis and treatment of these defects.

Case reports

Case 1. L.A., a one month-old Negro boy, was admitted to Babies Hospital for cardiac evaluation. A murmur had been noted on the first day of life

and congestive heart failure and pneumonia occurred at nine days of age. The infant was digitized and treated with penicillin and kanamycin at another hospital. Transfer to Babies Hospital was arranged when the baby did not improve.

Physical examination disclosed a vigorous active infant in moderate respiratory distress. The weight was 2.6 kilograms, heart rate 120, respiratory rate 100. The lungs were clear to auscultation. Examination of the heart revealed a grade III/VI blowing pansystolic murmur at the lower left sternal border. S₂ was single. The liver was palpable 3 cm below the right costal margin. The chest x-ray showed an enlarged heart with pulmonary vascular overcirculation. An electrocardiogram (ECG) showed a QRS axis of +120°, an RrS in V₁ and right ventricular predominance. Oxygen saturation in room air was 90 per cent. The infant showed no improvement on a maximal cardiotoxic regimen. Three days following admission cardiac catheterization and selective right ventricular angiography demonstrated total anomalous pulmonary venous drainage via the coronary sinus. Left ventricular angiocardiology was not performed. The data are summarized in Table 1.

Three days following the cardiovascular study the total anomalous venous drainage was totally repaired by means of cardiopulmonary bypass. The postoperative course was characterized by recurrent pneumonia and atelectasis with Pco₂ ranging from 40.9 to 61 mm Hg. Persistent pulmonary failure led to long term administration of high oxygen concentrations. A lower left sternal border holosystolic murmur persisted. Two months postoper-

From the Division of Pediatric Cardiology, Department of Pediatrics, and the Department of Radiology, College of Physicians and Surgeons, Columbia University and Babies Hospital, Columbia Presbyterian Medical Center, New York, N.Y.

Title was supported by Training Grant HE05389-12 of the National Institutes of Health.

Received for publication May 13, 1972.

Reprints: Carl N. Steeg, MD, Columbia University College of Physicians and Surgeons, Medical Center 622 W 168 St., New York, N.Y. 10032.

specifically discussed with a view to early diagnosis and therapy. The relevant literature is reviewed.

The authors wish to thank Mrs Mickey Mitchell for her able assistance in the preparation of this manuscript.

REFERENCES

1. Horn H and Saphir O. The involvement of the myocardium in tuberculosis: a review of the literature and report of three cases. *Am Rev Tuberc* 32:492 1935.
2. Auerbach O and Guggenheim A. Tuberculosis of the myocardium: a review of the literature and a report of six new cases. *Q Bull Sea View Hosp* 2:264 1937.
3. Kinare S G and Deshmukh M M. Complete atrioventricular block due to myocardial tuberculosis: report of a case. *Arch Pathol* 88:684 1969.
4. Pomerance A. Tuberculoma of the interventricular septum. *Br Heart J* 25:412 1963.
5. Rawls W J, Shuford W H, Logan W D, Hurst J W and Schlant R C. Right ventricular outflow tract obstruction produced by a myocardial abscess in a patient with tuberculosis. *Am J Cardiol* 21:738 1968.
6. Jones K P and Tilden I L. Tuberculous myocardial aneurysm with rupture and sudden death from tamponade. *Hawaii Med J* 1:295 1942.
7. Campos R, De C J, and Maranzana L F. Tuberculosis del corazón: presentación de un caso. *Rev Tuberc Lima* 10:12 1950.
8. Townsend R. Contributions to pathological anatomy. Case V. Death from asphyxia caused by large tuberculous masses developed in the parietes of the left auricle compressing the trunks of the pulmonary veins so as to reduce their diameter to that of a crow quill thereby reducing the return of blood from the lungs. *Dublin J Med Sci* 1:176 1832.
9. Adamson W W. A case of tuberculosis of the myocardium. *J Pathol Bacteriol* 23:399 1970.
10. Anders J M. Tuberculosis of the myocardium. *J A M A* 39:1081 1902.
11. Gouley B A, Bellet S and McMillan T M. Tuberculosis of the myocardium. Report of six cases with observations on involvement of coronary arteries. *Arch Intern Med* 51:244 1933.
12. James T N. Myocardial infarction and atrial arrhythmias. *Circulation* 24:761 1961.
13. Weigert K. Zur Lehre von der tuberculose und von verwandten Erkrankungen. *Virchows Arch (Pathol Anat)* 77:269 1879.
14. Ziegler E. *Lehrbuch allgem u speciel Pathol u patholog Anat*. G Fischer, Jena 1892. Quoted by Horn H, and Saphir O.
15. Rauchwerger S M and Rogers R J. Tuberculoma of the myocardium. A case report. *Am Heart J* 34:280 1947.
16. Banyu A L and Van Hecke L J. Tuberculoma of the myocardium: report of case. *Wisconsin Med J* 36:721 1937.
17. *Mitrocorditus A Pneumaticum Instrumentum Circulandi Sanguinis*. Frankfurt 1665. T M Gotzau chap 10 p 88.
18. Raviart G. La tuberculose du myocarde. *Arch Med Exp Anat Pathol* 11:141 1906.
19. Fontoynot M. Tuberculose du myocarde. *Bull Soc Anal Paris* 20:101 1897.

were heard over both posterior lung fields. Cardiac examination revealed a prominent systolic thrill at the left lower sternal border. The second sound was single. A grade IV/VI rough holosystolic murmur was heard along the left lower sternal border. The liver was palpable 7 cm below the right costal margin. There was no cyanosis. The peripheral pulses were strong and equal. Chest x ray showed an enlarged heart with marked increase in prominence of the pulmonary vasculature (Fig 1). The ECG revealed the QRS to be $+110^\circ$ with right ventricular preponderance (Fig 2). P waves in Lead II were 2.5 mm and peaked.

The infant improved following treatment with digoxin, furosemide and oxygen. One week following admission cardiac catheterization and right ventricular angiocardiography documented total anomalous pulmonary venous drainage to the right atrium (apparently via the coronary sinus) (Fig 3). The catheterization data showed right ventricular hypertension and some right ventricular outflow tract obstruction within a 52 mm peak systolic gradient between the pulmonary artery and the body of the right ventricle (Table 1).

A selective left ventricular angiogram was also performed which showed a large left-to-right shunt at the ventricular level (Fig 4).

The infant's condition remained stable and he was discharged on digoxin three days following the procedure. He is presently doing well at home at 18 months of age.

Discussion

The reported incidence of total anomalous pulmonary venous drainage with a ventricular septal defect as the sole complicating lesion varies. Brody⁴ reviewing 102 cases of anomalous pulmonary venous drainage of which 37 were total, found no case of associated ventricular septal defect. Burroughs and Edwards⁵ did not find a ventricular septal defect as a single associated malformation in their review of 188 cases of total anomalous pulmonary venous drainage although 66 had other multiple major cardiac anomalies including in many cases the asplenia syndrome. In a review of 30 patients with total anomalous pulmonary venous drainage Gott and associates⁶ found six cases to have associated anomalies but none was an isolated ventricular septal defect. The extensive review of Nakib and colleagues⁷ and Blake and colleagues⁸ also failed to show ventricular septal defect alone as a frequent complicating lesion.

On the other hand Bonham Carter and associates¹ in an analysis of 75 children with total anomalous pulmonary venous drainage found that of 17 with major associated anomalies four had ventricular



Fig 1 Admission chest roentgenogram (Case 3) showing cardiomegaly with increased vascularity.

septal defect. Our experience since 1966 consists of 19 patients with total anomalous pulmonary venous drainage, three of whom had ventricular septal defects.

When total anomalous pulmonary venous drainage with pulmonary hypertension exists as an isolated lesion, we have reported in a previous publication that early surgical correction of the defect by means of cardio-pulmonary bypass has been a successful method of treatment at this institution.⁹ However, when a complicating ventricular septal defect is present, this may not be the best approach to management. Indeed, failure to diagnose the additional anomaly may prove fatal if correction of the total anomalous venous drainage alone is undertaken.

In all three of these cases the diagnosis of ventricular septal defect with left-to-right shunt was considered on the basis of the clinical presentation including especially the auscultatory findings. In two of these cases, however, after the documentation of total anomalous pulmonary venous drainage during the initial catheterization, the possibility of ventricular septal defect was not further investigated—partly in view of the supposed rarity of this association. The murmurs were then attributed to tricuspid insufficiency. In retrospect, however, the holosystolic left sternal border murmurs were not caused by tricuspid insufficiency but were secondary to the ventricular septal defects as initially suspected. In one of these cases, repeat catheterization did reveal the presence of the

Table I Cardiac catheterization data

| Patient | SVC | | RA | | RV | | PA | | LA | | L3 | | PVR (RU/M ²) | |
|---------|-------|-----|-----|------|-----|----------|-----|--------|-----|----|-----|----|-----------------------------|-----|
| | Sat * | P* | Sat | P | Sat | P | Sat | P | Sat | P | Sat | P | | |
| I. A | | | | | | | | | | | | | | |
| No 1 | 81 | | 85 | 1M-6 | 80 | 65/12ED† | 88 | 65/24 | 45 | 95 | M-3 | 96 | 4/°ED | 3.6 |
| No 2 | 72 | M-2 | 73 | M-2 | 82 | 33/3ED | 82 | 35/18 | 28 | | | | | 2.2 |
| N G | 80 | | | M-5 | 93 | 102/7ED | 87 | 85/45 | 68 | 85 | M-7 | 83 | 110/10ED | 4.4 |
| R R | 84½ | | | | | | | | | | | | | |
| | 69 | | 97 | M-7 | 89 | 86/6½ED | 91 | 3°/10½ | 20 | | M-7 | 88 | 88/- | 0.9 |

*Sat = Oxygen saturation P = pressure

†M = mean

‡ED = End diastolic

||VC

||Not simultaneous

||Direct withdrawal

actively a second catheterization was carried out and a left-to-right shunt at the ventricular level was suspected on the basis of oxygen saturation data. The pulmonary artery pressure was only minimally increased (Table I) and pulmonary vascular resistance (PVR) was calculated at 2.2 resistance units (RU).

The infant continued to do poorly over the succeeding months. Feedings were not taken well and respiratory distress continued. Despite the use of a feeding gastrostomy the infant failed to thrive. His physical status gradually deteriorated and he died 7½ months postoperatively.

Postmortem examination. The heart and lungs weighed 250 grams (normal = 119 ± 10). The total anomalous pulmonary venous drainage was well repaired and there was no visible obstruction to pulmonary venous return. A ventricular septal defect 0.8 cm in diameter was found above the crista supraventricularis. The right ventricle was dilated as was the main pulmonary artery, which was larger in diameter than the ascending aorta. The left ventricle was slightly larger than the right.

Microscopic examination of the lungs revealed hypertrophy of arteries and veins with mild muscular arterial hyperplasia (Stage II Pulmonary Vascular Disease)¹² as well as septal and focal interlobular fibrosis. These fibrotic changes may represent the effects of prolonged oxygen administration.

Case 2 N G. A 4-month-old Caucasian girl was admitted to Babies Hospital for cardiac surgery. The history was unremarkable until 2½ months of age when tachypnea and increasing fatigability were noted. A murmur compatible with the clinical diagnosis of ventricular septal defect was heard one month later.

The infant was admitted to another hospital and digitalized with minimal improvement. Cardiac catheterization and angiography showed total anomalous pulmonary venous drainage via a left superior vena cava. A left ventricular angiogram was not done. She was then transferred to Babies Hospital.

On admission the infant was noted to be apyretic and in mild respiratory distress. The weight was 4.1 kilograms, heart rate 150, respiratory rate 70. The lungs were clear to auscultation. Examination of the heart revealed a visible right ventricular impulse, a grade III/VI holosystolic murmur at the left sternal border and a single second sound. There was no hepatomegaly or peripheral edema.

The ECG showed an axis of +135 with right ventricular enlargement. The T waves were upright in V₁. A chest x-ray showed cardiomegaly with a wide superior mediastinum and pulmonary vascular engorgement.

Because of the murmur suggestive of ventricular septal defect the infant was recatheterized. The catheterization data confirmed total anomalous pulmonary venous drainage via the superior vena cava with pulmonary hypertension (Table I). An angiocardiogram performed with the catheter in the left ventricle demonstrated the presence of a large high ventricular septal defect with left-to-right shunt. Two days later an open heart operation was performed in an attempt at total correction. In addition to the total anomalous pulmonary venous drainage to the left innominate vein a large membranous ventricular septal defect was found. Correction of these anomalies required 80 minutes of cardiopulmonary bypass. Four-hour postoperatively the patient became severely acidotic. Multiple cardiac arrests ensued and she died eight hours postoperatively. An autopsy was not obtained.

Case 3 R R. A 2-month-old Caucasian boy was admitted to Babies Hospital because of congestive heart failure. He was the product of an uncomplicated term pregnancy and appeared normal at birth. A heart murmur was noted at one week of age but the infant thrived at home with good weight gain. One week prior to admission tachypnea, fatigue with feedings and excessive sweating were noted.

Physical examination revealed a well-developed, well-nourished, acyanotic infant in moderate respiratory distress. The weight was 4.6 kilograms, heart rate 170, respiratory rate 90. Bilateral moist rales



Fig 4 Frontal (left) and lateral (right) angiocardiograms (Case 3) during injection of radiopaque contrast material into the relatively small left ventricle. A substantial left-to-right shunt via a ventricular septal defect opacifies the right ventricle and pulmonary arteries. The right ventricular outflow tract gradient was identified by catheter withdrawal.

lar stroke output tends to increase blood flow in the pulmonary circuit and at the same time reduce the systemic output. Such hemodynamics require relatively large flows from right to left across the interatrial septum. If resistance in the pulmonary circuit is high, left-to-right shunting via the ventricular septal defect would be limited, and in rare instances it might be speculated that right-to-left shunting through such a ventricular septal defect would occur, reducing pulmonary blood flow while augmenting systemic output.

In the absence of a ventricular septal defect, surgical elimination of the large pulmonary vein-right atrial shunt reduces right ventricular return and right ventricular output. The pulmonary artery pressure is correspondingly reduced even assuming no change in pulmonary vascular resistance. When a large ventricular septal defect is present, however, the reduced right ventricular output would tend to be matched by a corresponding increase in the left-to-right shunt via the ventricular septal defect. Thus, by repair of the anomalous venous connections alone, the work of pumping a large left-to-right shunt is shifted from the right to the left ventricle. This radical change in the output requirements for the left ventricle may not be tolerated. In Case 1, the limited size of the ventricular septal defect probably prevented immediate catastrophe following repair of the total anomalous pulmonary venous drainage, and death seemed unrelated to the interventricular communication.

Since the ventricular septum is usually intact in total anomalous pulmonary venous drainage, palliative pulmonary artery banding has not been considered. However, when a large ventricular septal defect complicates total anomalous pulmonary venous drainage in a critically ill infant with pulmonary hypertension and congestive heart failure, a palliative operation—banding of the pulmonary artery—might be considered. This procedure would be expected to reduce both pulmonary blood flow and pulmonary hypertension while simultaneously augmenting systemic output. Such beneficial effects are suggested by the favorable course of R.R. (Case 3), who had a moderate obstruction and pressure gradient across the pulmonary outflow tract.

Summary

Three patients with total anomalous pulmonary venous drainage and ventricular septal defect with left-to-right shunt are presented. The ventricular septal defect was diagnosed with left ventricular angiography in two patients and confirmed on postmortem examination in the third.

The complex hemodynamic implications of this combination of lesions are discussed. Pulmonary artery banding is suggested as a possible method of management in early infancy.

When total anomalous pulmonary venous drainage is demonstrated in infants initially suspected of having large ventricular septal defects on the basis of auscultation, left ventricular angiography is mandatory. Preoperative diagnosis of a compli-

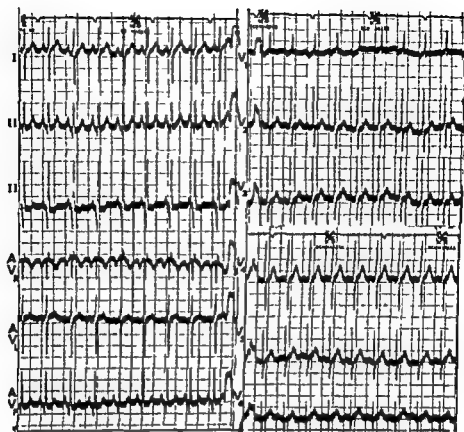


Fig 2 Admission ECG (Case 3) Axis $+110^\circ$ Right ventricular preponderance P waves are peaked in Lead II



Fig 3 Frontal angiogram (Case 3) showing late phase of pulmonary vein opacification 3 sec following injection of radiopaque contrast material into right ventricle. The expected opacification of the left atrium is not seen. Instead the pulmonary veins opacify the right atrium (RA) apparently via the coronary sinus (arrows)

complicating ventricular septal defect prior to the attempt at corrective surgery

In the third case, the initial diagnosis of ventricular septal defect was confirmed by a left ventricular angiogram in the early phases of the catheterization, and total anomalous pulmonary venous drain-

age was not really suspected until subsequent completion of the right sided study

Since left and right ventricular oxygen saturations are similar in total anomalous pulmonary venous drainage oxygen saturation determinations cannot be expected to be of great value in the detection of a complicating ventricular septal defect. Oxygen saturation data however might have been of value in one of our patients (N G Case 2), in whom a consistent 'step down' in saturation was noted on entering the pulmonary artery from the right ventricle (Table I). In this instance left heart saturations were somewhat lower than those in the right ventricle—presumably due to streaming of less saturated inferior vena caval blood into the left heart via the foramen ovale. Thus a left to right shunt at the ventricular level may have been paradoxically responsible for the modest fall or 'step down' of oxygen saturation in the pulmonary artery.

The hemodynamic significance of a large ventricular septal defect in total anomalous pulmonary venous drainage is complex. In our cases left to right shunts occurred via the ventricular septal defect. This diversion of some of the left ventricu-

Experimental and laboratory reports

Early arrhythmias following experimental coronary occlusion in conscious dogs and their modification by beta-adrenoceptor blocking drugs*

Maidul I Khan MB BS PhD FRCP(C)**

John T Hamilton BSc PhD (Edin)

George W Manning MD PhD FRCP(C) FACC

London Ontario, Canada

The influence of the beta adrenoceptor blocking agents in experimental coronary occlusion has been studied by many investigators¹⁻⁸ but all of their experiments were performed either in anesthetized open chest animals or in conscious animals one to five days after coronary occlusion. As early as 1939 it was pointed out that general anesthesia by itself significantly reduces the incidence of ventricular arrhythmias following the ligation of either branch of the left coronary artery.⁹ It has also been shown that the hemodynamic effects of the beta adrenoceptor blocking drugs differ considerably between conscious and anesthetized animals.¹⁰ Harris and associates¹¹ described three phases following abrupt coronary occlusion in anesthetized dogs: first the period of marked danger from ventricular ectopic beats and fibrillation usually lasting for 10 to 15 minutes from the time of occlusion; the second

period with a decline of ectopic activity and lasting until the development of ventricular tachycardia—between 45 and 8 hours; and a third phase characterized by delayed arrhythmias lasting for two to five days. Those studies done on the second or third day of coronary occlusion thus missed the phase where the danger of ventricular ectopic beats, fibrillation and sudden death is greatest.

The present investigation was designed to evaluate the influence of a group of beta adrenoceptor blocking drugs on the arrhythmias that complicate the early phase of acute coronary occlusion in conscious dogs. The particular drugs selected for study permitted a definition of the relative roles played by generalized beta adrenoceptor blockade by cardioselective beta blockade and by the local anesthetic properties of these substances. Their effect on heart rate, blood pressure

*In the Department of Pharmacology, Medical Sciences Centre, University of Western Ontario and the Cardiac Sciences Unit, Hammersmith Hospital, London, Ontario, Canada.

This study was supported by grants from the Ontario Heart Foundation.

Received for publication May 4, 1972.

Reprint requests to Dr. J. T. Hamilton, Department of Pharmacology, University of Western Ontario, London, Ontario, Canada.

These studies were presented at the meeting of the Royal College of Physicians and Surgeons of Canada, June 29-30, 1972, at the Cardiac Society of the United Kingdom, June 1970, both in Montreal and at the International Symposium on Atherosclerosis, June 1971, at Toronto, Canada.

**On leave from Foundation F. B. B.

cating ventricular septal defect is essential to optimal management

REFERENCES

- 1 Bonham Carter R E, Capriles M and Noe Y Total anomalous pulmonary venous drainage a clinical and anatomical study of 75 children *Br Heart J* 31:45 1969
- 2 Darling R C Rothney W B and Craig J M Total pulmonary venous drainage into the right side of the heart *Lab Invest* 6:44 1957
- 3 Heath D and Edwards J E The pathology of hypertensive pulmonary vascular disease. A description of six grades of structural changes in the pulmonary arteries with special reference to congenital cardiac septal defects *Circulation* 18:533 1958
- 4 Brody H Drainage of the pulmonary veins into the right side of the heart *Arch Pathol* 33:221, 1942
- 5 Burroughs J T and Edwards J E Total anomalous pulmonary venous connection *Am Heart J* 59:913 1960
- 6 Gott V L Lester R G Lillehei C W and Vinco H L Total anomalous pulmonary return. An analysis of 30 cases *Circulation* 13:543 1956
- 7 Nakib A Moller J H Kanjuh V I and Edwards J F Anomalies of the pulmonary veins *Am J Cardiol* 20:77 1967
- 8 Blake H A Hall R J and Manion W C Anomalous pulmonary venous return *Circulation* 32:406 1965
- 9 Gersony W M Bowman F O Jr Steeg C N Hayes C J Jesse M J and Malm J R Management of total anomalous pulmonary venous drainage in early infancy *Circulation* 43 (Suppl 1) 19 1971

Experimental and laboratory reports

Early arrhythmias following experimental coronary occlusion in conscious dogs and their modification by beta-adrenoceptor blocking drugs*

Maidul I Khan MB BS PhD FRCP(C)*

John T Hamilton BSc PhD (Edin)

George W Manning MD PhD FRCP(C) FACC

London Ontario Canada

The influence of the beta adrenoceptor blocking agents in experimental coronary occlusion has been studied by many investigators¹⁻⁴ but all of their experiments were performed either in anesthetized open chest animals or in conscious animals one to five days after coronary occlusion. As early as 1939 it was pointed out that general anesthesia by itself significantly reduces the incidence of ventricular arrhythmias following the ligation of either branch of the left coronary artery.⁵ It has also been shown that the hemodynamic effects of the beta adrenoceptor blocking drugs differ considerably between conscious and anesthetized animals.⁶⁻⁸ Harris and associates¹¹ described three phases following abrupt coronary occlusion in anesthetized dogs: first the period of marked danger from ventricular ectopic beats and fibrillation usually lasting for 10 to 15 minutes from the time of occlusion the second

period with a decline of ectopic activity and lasting until the development of ventricular tachycardia—between 45 and 8 hours and a third phase characterized by delayed arrhythmias lasting for two to five days. Those studies done on the second or third day of coronary occlusion thus missed the phase where the danger of ventricular ectopic beats fibrillation and sudden death is greatest.

The present investigation was designed to evaluate the influence of a group of beta adrenoceptor blocking drugs on the arrhythmias that complicate the early phase of acute coronary occlusion in conscious dogs. The particular drugs selected for study permitted a definition of the relative roles played by generalized beta adrenoceptor blockade by cardio selective beta blockade and by the local anesthetic properties of these substances. Their effect on heart rate, blood pressure

*From the Department of Pharmacology, Medical Sciences Centre, University of Western Ontario and the Cardiovascular Society, Hillier, London, Ontario, Canada.

Received for publication September 1, 1972; accepted for publication October 1, 1972.

Reprint requests to Dr. J. T. Hamilton, Department of Pharmacology, University of Western Ontario, London, Ontario, Canada.

These studies were presented at the meeting of the Royal College of Physicians and Surgeons of Canada in June 1970 both at Montreal and at the 11th Symposium on Arrhythmias, June 1972, at Toronto, Canada.

*Ontario Hospital Foundation Fellow.

Table 1 The effect of various drugs compared to normal saline injection on the mean heart rate of conscious dogs

| Mean heart rate \pm standard error of the mean | | | | |
|---|-----------------|-----------------|-----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 121.9 \pm 5.0 | 123.2 \pm 5.0 | +1.3 \pm 2.2 | > 0.5 |
| 2 dl propranolol 0.1 mg/kg | 107.4 \pm 4.8 | 99.6 \pm 4.6 | -7.8 \pm 2.8 | < 0.01 |
| 3 dl propranolol 1.0 mg/kg | 122.5 \pm 7.3 | 110.9 \pm 5.6 | -11.6 \pm 3.8 | < 0.01 |
| 4 MJ 1999 0.2 mg/kg | 118.7 \pm 2.7 | 112.8 \pm 2.8 | -6.0 \pm 2.0 | < 0.01 |
| 5 MJ 1999 3.2 mg/kg | 95.4 \pm 6.5 | 89.6 \pm 4.4 | -5.8 \pm 5.3 | > 0.3 |
| 6 d propranolol 0.07 mg/kg | 117.3 \pm 9.5 | 112.3 \pm 9.5 | -5.1 \pm 5.1 | > 0.3 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 114.4 \pm 5.0 | 111.0 \pm 4.3 | -3.4 \pm 3.1 | > 0.2 |
| 8 AY 21 011 1.4 mg/kg | 109.6 \pm 7.1 | 105.8 \pm 7.2 | -3.8 \pm 2.6 | > 0.1 |

and electrocardiogram were measured. The mortality rates found with these agents in experimental coronary occlusion in conscious dogs are published elsewhere.¹²

Methods and materials

Adult mongrel dogs (5 to 18 kilograms) were anesthetized with pentobarbital (25 to 30 mg per kilogram of body weight) and a loose ligature was placed around the circumflex branch of the left coronary artery at its origin under direct vision as described previously.¹³ The ends of the ligature were exteriorized the chest was closed in layers, and the animal was allowed to recover for 48 to 72 hours. Then the right femoral artery and vein were cannulated under local anesthesia. All drugs were administered through the venous catheter. Direct arterial pressure (Statham P23 db transducer) and electrocardiogram (usually Lead II from subcutaneous needle electrodes) were recorded simultaneously on a Sanborn (Model 67 1600) multi-channel direct writer. Magnetic tape recordings (Sony) made at the same time were used for subsequent detailed analysis of arrhythmias.

Control animals received normal saline and treated animals received an appropriate amount of the assigned drug dissolved in normal saline in exactly the same manner. All doses are expressed in terms of the salt used. Five minutes later the exterior ends of the ligature were pulled to totally occlude the artery. Blood pressure and ECG were recorded continuously for the

first 10 minutes after coronary occlusions and every 15 minutes thereafter for a period of two hours or until death. Continuous tape recordings of ECG were made throughout the two hour period of observation in the first 65 experiments, but subsequently, this was reduced to one hour.

A total of 141 dogs was randomly allocated as follows:

- Group 1 normal saline 2 to 3 ml (n = 25)
- Group 2 dl propranolol (Inderal) * 0.1 mg per kilogram of body weight (n = 25)
- Group 3 dl propranolol * 1.0 mg per kilogram of body weight (n = 10)
- Group 4 MJ 1999 (Sotalol) † 0.2 mg per kilogram of body weight (n = 25)
- Group 5 MJ 1999 † 3.2 mg per kilogram of body weight (n = 12)
- Group 6 d propranolol * 0.07 mg per kilogram of body weight (n = 16)
- Group 7 d propranolol 0.07 mg per kilogram of body weight + MJ 1999 0.2 mg per kilogram of body weight (n = 12)
- Group 8 AY 21 011 (Prietolol) * 1.4 mg per kilogram of body weight (n = 16)

* Vicer Laboratories, Montreal, Canada.
† Mead Johnson & Co., Evansville, Ind.

The doses of the drugs were selected from the results of preliminary experiments in conscious dogs in which their ability to antagonize the chronotropic effect of 140 protorenol (1.5 µg per kilogram of body weight) was determined. The doses used in Groups 2, 4, and 8 were equiaffective and caused 50 per cent blockade of isoproterenol induced tachycardia and those used in Groups 3 and 5 caused nearly 100 per cent blockade. The dose of *d* propranolol used in Groups 6 and 7 has been shown to have an equal local anesthetic effect to that of the *dl* propranolol used in Group 2.¹²

For detailed analysis of arrhythmias the magnetic tape was played back using a Philips tape recorder with attached arrangement for visual display of electrocardiogram on a Dumont 401B oscilloscope. A Hewlett Packard electrocardiogram machine in the system allowed direct write-out of any segment of taped electrocardiogram and a Philips Analog 7 recorder was used for rerecording and playbacks at slower than recorded rate when required.

The significance of the effects on heart rate, blood pressure and PR interval was tested by Student's *t* test using paired data and the mean changes were subjected to overall analysis of variance. The immediate effects of coronary occlusion on the heart rate and blood pressure were analyzed in the same way. The Chi square method was used to test the significance of the difference in the number of experimental animals exhibiting a particular cardiac arrhythmia. For numerical analysis the ventricular ectopic beats were expressed in terms of their frequency per minute in each dog during the three 5 minute periods following coronary occlusion. The difference between each of the parameters tested was considered significant only when *p* values were less than 0.05.

Results

The effects of beta-adrenoceptor blocking drugs in conscious dogs. No unusual general effects—e.g. vomiting, diarrhea, excitement, tremor, convulsions, stupor or shortness of breath—were noted following the administration of the drugs used in this study.

There was a decrease in the mean heart rate (Table I) and an increase in both systolic and diastolic blood pressures (Ta-

ble II) following the administration of the beta adrenoceptor blocking drugs. When the changes induced by normal saline were compared with the changes caused by all the treatments, taken as a group, the difference was significant ($p < 0.05$). Except for AY 21 011, all beta adrenoceptor blocking drugs in the doses used caused a significant prolongation of the PR interval (Table III).

The effects of sudden coronary occlusion in conscious dogs. There was a significant increase in heart rate in all groups immediately following the coronary occlusion, reaching a peak between 30 and 120 seconds, then gradually returning towards the pre-occlusion level (Fig. 1).

There was a decrease in both systolic and diastolic blood pressures in all groups immediately following the coronary occlusion, reaching its lowest level within two minutes and gradually coming back towards but not reaching the pre-occlusion level within the 2 hour period (Fig. 2A and B).

Systolic pressures decreased more than diastolic so that the pulse pressure following coronary occlusion remained below the pre-occlusion level during the period of observation. For the systolic blood pressure the difference between the pre-occlusion level and the maximal drop following coronary occlusion analyzed as paired data was significant in all groups ($p < 0.02$ to 0.001) except in the group treated with *d* propranolol ($p > 0.05$) but there were no significant differences among the various treatment groups. The maximal fall in diastolic blood pressure following coronary occlusion was statistically significant in Groups 1, 2, and 7 but again was not significantly different among these groups.

Cardiac arrhythmias. The incidence of ventricular fibrillation (VF) in Groups 2, 4, 5, 7, and 8 was significantly lower than that in Groups 1 and 6 (Table IV). The 40 per cent incidence of VF in Group 3 when contrasted with the 72 per cent in the control group approached significance.

Ventricular ectopic beats were classified into (1) ventricular premature beats (PVB) when they appeared singly or in groups of three or less, and (2) ventricular tachycardia (VT) when they occurred in groups of four or more. When present they usually appeared within seconds of coro-

Table 11A The effect of various drugs compared to normal saline on the blood pressure in conscious dogs

| Mean systolic blood pressure (mm Hg) \pm standard error of the mean | | | | |
|---|-----------------|-----------------|----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 153.4 \pm 4.8 | 154.3 \pm 4.3 | +0.9 \pm 2.0 | > 0.5 |
| 2 dl propranolol 0.1 mg/kg | 154.9 \pm 4.0 | 164.8 \pm 4.2 | +9.9 \pm 1.7 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 155.0 \pm 6.7 | 161.4 \pm 5.6 | +6.4 \pm 2.4 | < 0.05 |
| 4 MJ 1999 0.2 mg/kg | 147.1 \pm 3.7 | 151.8 \pm 3.9 | +4.6 \pm 1.9 | < 0.05 |
| 5 MJ 1999 3.2 mg/kg | 158.5 \pm 6.0 | 161.0 \pm 7.4 | +2.5 \pm 4.7 | > 0.5 |
| 6 d propranolol 0.7 mg/kg | 164.0 \pm 6.7 | 168.5 \pm 8.1 | +4.5 \pm 3.3 | > 0.1 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 164.3 \pm 5.5 | 170.0 \pm 6.8 | +5.7 \pm 4.4 | > 0.1 |
| 8 Ay 21 011 1.4 mg/kg | 162.1 \pm 5.6 | 168.6 \pm 5.8 | +6.5 \pm 3.4 | > 0.1 |

Table 11B The effect of various drugs compared to normal saline on the blood pressure in conscious dogs

| Mean diastolic blood pressure (mm Hg) \pm standard error of the mean | | | | |
|--|----------------|-----------------|-----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 91.0 \pm 3.2 | 92.0 \pm 3.2 | +1.0 \pm 2.0 | > 0.5 |
| 2 dl propranolol 0.1 mg/kg | 92.6 \pm 2.9 | 102.7 \pm 3.3 | +10.1 \pm 1.8 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 91.0 \pm 3.1 | 97.4 \pm 3.8 | +6.4 \pm 1.5 | < 0.01 |
| 4 MJ 1999 0.2 mg/kg | 83.4 \pm 2.6 | 89.4 \pm 2.8 | +6.0 \pm 1.6 | < 0.001 |
| 5 MJ 1999 3.2 mg/kg | 86.3 \pm 3.0 | 92.7 \pm 5.0 | +6.3 \pm 4.5 | > 0.1 |
| 6 d propranolol 0.07 mg/kg | 95.5 \pm 4.5 | 95.8 \pm 3.9 | +0.3 \pm 2.2 | > 0.5 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 95.8 \pm 4.2 | 106.2 \pm 5.7 | +10.3 \pm 6.6 | > 0.1 |
| 8 Ay 21 011 1.4 mg/kg | 89.5 \pm 3.4 | 97.1 \pm 3.3 | +7.6 \pm 3.1 | < 0.05 |

Table 111 The effect of various drugs compared to normal saline on the mean PR interval in conscious dogs

| Mean PR interval (msec) \pm standard error of the mean | | | | |
|--|-----------------|-----------------|-----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 94.7 \pm 2.5 | 97.8 \pm 3.0 | +2.9 \pm 1.7 | > 0.05 |
| 2 dl propranolol 0.1 mg/kg | 110.0 \pm 4.4 | 121.3 \pm 4.4 | +11.3 \pm 1.4 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 102.5 \pm 3.0 | 113.9 \pm 3.6 | +11.4 \pm 1.7 | < 0.001 |
| 4 MJ 1999 0.2 mg/kg | 95.4 \pm 2.8 | 105.1 \pm 3.4 | +9.6 \pm 1.4 | < 0.001 |
| 5 MJ 1999 3.2 mg/kg | 110.6 \pm 7.1 | 121.5 \pm 6.1 | +10.9 \pm 2.7 | < 0.01 |
| 6 d propranolol 0.07 mg/kg | 102.4 \pm 5.5 | 105.7 \pm 5.9 | +3.3 \pm 1.6 | < 0.05 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 95.7 \pm 3.8 | 106.5 \pm 4.6 | +10.7 \pm 1.8 | < 0.001 |
| 8 Ay 21 011 1.4 mg/kg | 102.4 \pm 6.1 | 103.9 \pm 5.4 | +1.5 \pm 1.7 | > 0.3 |

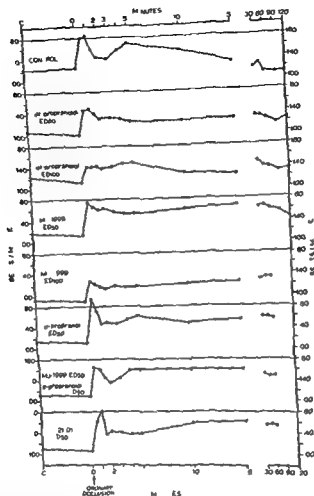


Fig. 1 Time-course of the mean heart rate in various groups. C = baseline value. O = coronary occlusion. C to O = effect of normal saline or drug. Minutes indicate time from coronary occlusion.

Table IV The incidence of ventricular fibrillation following sudden occlusion of the circumflex branch of the left coronary artery in conscious dogs

| Group | Number of dogs | Ventricular fibrillation | | Significance of the difference control vs treatment |
|--|----------------|--------------------------|----------|---|
| | | Number | Per cent | |
| 1 Control | 25 | 18 | 72 | |
| 2 dl propranolol 0.1 mg/kg | 25 | 5 | 20 | $p < 0.001$ |
| 3 dl propranolol 1.0 mg/kg | 10 | 4 | 40 | $0.05 < p < 0.1$ |
| 4 dl 1999 0.2 mg/kg | 25 | 7 | 28 | $p < 0.005$ |
| 5 dl 1999 3.2 mg/kg | 12 | 2 | 17 | $p < 0.005$ |
| 6 dl 1999 0.2 mg/kg + dl 1999 0.07 mg/kg | 16 | 11 | 69 | $\equiv > 0.75$ |
| 7 dl 1999 0.2 mg/kg | 12 | 2 | 17 | $p < 0.005$ |
| 8 dl 1999 1.4 mg/kg | 16 | 3 | 19 | $p < 0.001$ |

The difference in the incidence of ventricular fibrillation between the groups is highly significant ($p < 0.001$) by a chi-square test.

Table II A The effect of various drugs compared to normal saline on the blood pressure in conscious dogs

| Mean systolic blood pressure (mm Hg) \pm standard error of the mean | | | | |
|---|-----------------|-----------------|----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 153.4 \pm 4.8 | 154.3 \pm 4.3 | +0.9 \pm 2.0 | > 0.5 |
| 2 dl propranolol 0.1 mg/kg | 154.9 \pm 4.0 | 164.8 \pm 4.2 | +9.9 \pm 1.7 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 155.0 \pm 6.7 | 161.4 \pm 5.6 | +6.4 \pm 2.4 | < 0.05 |
| 4 MJ 1999 0.2 mg/kg | 147.1 \pm 3.7 | 151.8 \pm 3.9 | +4.6 \pm 1.9 | < 0.05 |
| 5 MJ 1999 3.2 mg/kg | 158.5 \pm 6.0 | 161.0 \pm 7.4 | +2.5 \pm 4.7 | > 0.5 |
| 6 d propranolol 0.7 mg/kg | 164.0 \pm 6.7 | 168.5 \pm 8.1 | +4.5 \pm 3.3 | > 0.1 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 164.3 \pm 5.5 | 170.0 \pm 6.8 | +5.7 \pm 4.4 | > 0.2 |
| 8 Ay 21 011 1.4 mg/kg | 162.1 \pm 5.6 | 168.6 \pm 5.8 | +6.5 \pm 3.4 | > 0.1 |

Table II B The effect of various drugs compared to normal saline on the blood pressure in conscious dogs

| Mean diastolic blood pressure (mm Hg) \pm standard error of the mean | | | | |
|--|----------------|-----------------|-----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 91.0 \pm 3.2 | 92.0 \pm 3.2 | +1.0 \pm 2.0 | > 0.5 |
| 2 dl propranolol 0.1 mg/kg | 92.6 \pm 2.9 | 102.7 \pm 3.3 | +10.1 \pm 1.8 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 91.0 \pm 3.1 | 97.4 \pm 3.8 | +6.4 \pm 1.5 | < 0.01 |
| 4 MJ 1999 0.2 mg/kg | 83.4 \pm 2.6 | 89.4 \pm 2.8 | +6.0 \pm 1.6 | < 0.001 |
| 5 MJ 1999 3.2 mg/kg | 86.3 \pm 3.0 | 92.7 \pm 5.0 | +6.3 \pm 4.5 | > 0.1 |
| 6 d propranolol 0.07 mg/kg | 95.5 \pm 4.5 | 95.8 \pm 3.9 | +0.3 \pm 2.2 | > 0.5 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 95.8 \pm 4.2 | 106.2 \pm 5.7 | +10.3 \pm 6.6 | > 0.1 |
| 8 Ay 21 011 1.4 mg/kg | 89.5 \pm 3.4 | 97.1 \pm 3.3 | +7.6 \pm 3.3 | < 0.05 |

Table III The effect of various drugs compared to normal saline on the mean PR interval in conscious dogs

| Mean PR interval (msec) \pm standard error of the mean | | | | |
|--|-----------------|-----------------|-----------------|---------------------------|
| Drug or saline | Before | After | Mean change | p value of the difference |
| 1 Normal saline | 94.7 \pm 2.5 | 97.8 \pm 3.0 | +2.9 \pm 1.7 | > 0.05 |
| 2 dl propranolol 0.1 mg/kg | 110.0 \pm 4.4 | 121.3 \pm 4.4 | +11.3 \pm 1.4 | < 0.001 |
| 3 dl propranolol 1.0 mg/kg | 102.5 \pm 3.0 | 113.9 \pm 3.6 | +11.4 \pm 1.7 | < 0.001 |
| 4 MJ 1999 0.2 mg/kg | 95.4 \pm 2.8 | 105.4 \pm 3.4 | +10.0 \pm 1.4 | < 0.001 |
| 5 MJ 1999 3.2 mg/kg | 110.6 \pm 7.1 | 121.5 \pm 6.1 | +10.9 \pm 2.7 | < 0.01 |
| 6 d propranolol 0.07 mg/kg | 102.4 \pm 5.5 | 105.7 \pm 5.9 | +3.3 \pm 1.6 | < 0.05 |
| 7 MJ 1999 0.2 mg/kg + d propranolol 0.07 mg/kg | 95.7 \pm 3.8 | 106.5 \pm 4.8 | +10.7 \pm 1.8 | < 0.001 |
| 8 Ay 21 011 1.4 mg/kg | 102.4 \pm 6.1 | 103.9 \pm 5.4 | +1.5 \pm 1.7 | > 0.3 |

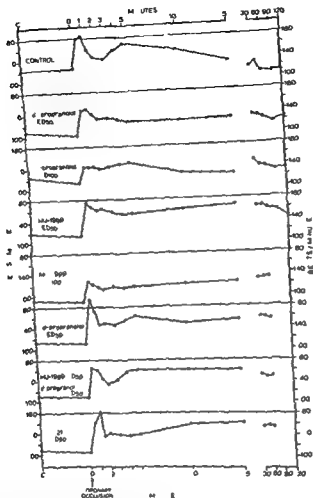


Fig. 1 Time course of the mean heart rate in various groups. C = base line value. O = coronary occlusion. C to O = effect of normal saline or drug. Minutes indicate time from coronary occlusion.

Table IV The incidence of ventricular fibrillation following sudden occlusion of the circumflex branch of the left coronary artery in conscious dogs

| Group | Number of dogs | Ventricular fibrillation | | Significance of the difference control vs treatment |
|--|----------------|--------------------------|----------|---|
| | | Number | Per cent | |
| 1 Control | 25 | 18 | 72 | |
| 2 d-propranolol 0.1 mg/kg | 25 | 5 | 20 | $p < 0.001$ |
| 3 d-propranolol 1.0 mg/kg | 10 | 4 | 40 | $0.05 < p < 0.1$ |
| 4 N1999 0.2 mg/kg | 25 | 7 | 28 | $p < 0.005$ |
| 5 N1999 3.2 mg/kg | 12 | 2 | 17 | $p < 0.005$ |
| 6 d-propranolol | 16 | 11 | 69 | $p > 0.75$ |
| 7 N1999 0.2 mg/kg + d-propranolol 0.07 mg/kg | 12 | 2 | 17 | $p < 0.005$ |
| 8 N1999 0.2 mg/kg + d-propranolol 0.07 mg/kg | 16 | 3 | 19 | $p < 0.001$ |

The difference in the incidence of ventricular fibrillation between the control group and the treated groups ($p < 0.001$) by one-way Chi-square test.

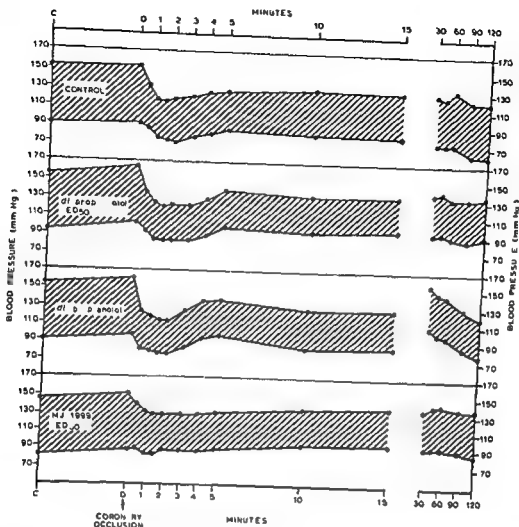


Fig. 2A Time course of the mean systolic and diastolic blood pressure in various groups. Abbreviations are the same as in Fig. 1.

nary occlusion rapidly increased in frequency reaching peaks at 2 to 5 minutes and then gradually decreased in frequency there being very few ectopic beats beyond 15 minute period following coronary occlusion (Fig. 3A and B). Ventricular ectopic beats were encountered in 69 to 92 per cent of dogs in the various groups (Table V). Ventricular tachycardia was noted in 12 to 72 per cent of animals in the various groups being lowest in Groups 2 and 5.

The total number of ectopic beats per dog per minute, without regard to type (all ectopic beats) during the first second and third 5 minute periods is shown in Table VI. When the incidence of all ectopic beats within the first five minutes in the control group is compared with all other groups taken together the difference is significant ($p < 0.05$) because of the low incidence in Groups 2, 5 and 6.

The number of PVB per minute during

the first, second and third 5 minute periods is shown in Table VII. The difference in their incidence between groups is not statistically significant in any of the three 5 minute periods.

When the incidence of ventricular tachycardia in the control group during the first 5 minute period is compared with that in all other groups taken together the difference is significant ($p < 0.01$) (Table VIII). The incidence of this arrhythmia in the second and third 5 minute periods was very low.

No dog in Groups 1, 4, 6 and 7 had atrioventricular block. Five out of ten dogs treated with the higher dose of *dl* propranolol developed progressive first, second and third degree atrioventricular block, three ending in cardiac standstill. Four out of sixteen dogs treated with AY 21 011 developed progressive heart block, two ending in cardiac standstill. Of the two

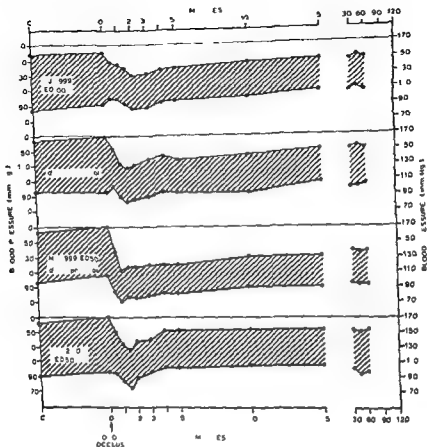


Fig. 2B Time course of the mean systolic and diastolic blood pressure in various groups. Abbreviations are the same as in Fig. 1.

dogs in the group treated with 0.1 mg per kilogram of body weight *dl* propranolol developed heart block; one ended in cardiac standstill. The incidence of severe heart block in Groups 3 and 8 is significantly higher than that in the control group ($p < 0.005$).

No dog in the control group showed sinus arrest but episodes of sinus arrest were noted in the remaining groups: one dog in Group 4, two in each of the Groups 2, 3, 5 and 6, three in Group 8 and four in Group 7.

Atrial tachycardia was noted in one dog in each of Groups 1, 2 and 6 and atrial fibrillation in one dog in Group 2.

Discussion

The effects of the drugs on the cardiovascular system are comparable to those in other studies^{2,14} but the extent of reduction in heart rate was not always as great. The

reduction in heart rate was statistically significant only with the lower dose of *MJ* 999 and the two doses of *dl* propranolol. Despite the fact that *AY* 21 011 possesses a mild intrinsic sympathomimetic effect and has been shown to increase the heart rate in anesthetized cats⁴ in most clinical studies it reduced the heart rate.^{15,16}

The rise of blood pressure after these drugs may be explained by an exaggerated alpha receptor response of the peripheral blood vessels to catecholamines.¹⁷ The fact that the larger doses of *dl* propranolol and *MJ* 999 caused less elevation than the lower doses is most likely due to a greater fall in cardiac output. The only discernible change in the electrocardiogram was a significant prolongation of the PR interval in all groups except the *AY* 21 011 (probably because of its intrinsic sympathomimetic action).¹⁸ These results are similar to those reported by others.^{10,21}

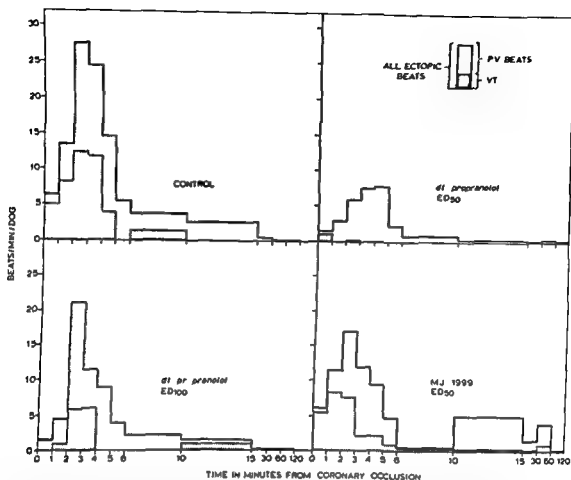


Fig 3A Ventricular ectopic beats following sudden coronary occlusion expressed in terms of beats/min/dog. Top time = all ectopic beats; hatched area = ectopic beats appearing in groups of four or more—ventricular tachycardia (VT); clear area = ectopic beats appearing singly or in groups of three or less—premature ventricular beats (PVB).

Table V Incidence of ventricular ectopic beats

| Group | Number of dogs | Premature ventricular beats | | Ventricular tachycardia | |
|---|----------------|-----------------------------|----------|-------------------------|----------|
| | | Number | Per cent | Number | Per cent |
| 1 Normal saline | 25 | 21 | 84 | 16 | 64 |
| 2 dl propranolol 0.1 mg/Kg | 25 | 17 | 68 | 3 | 12 |
| 3 dl propranolol 1.0 mg/Kg | 10 | 8 | 80 | 4 | 40 |
| 4 MJ 1999 0.2 mg/Kg | 25 | 23 | 92 | 18 | 72 |
| 5 MJ 1999 3.2 mg/Kg | 12 | 9 | 75 | 3 | 25 |
| 6 dl propranolol 0.07 mg/Kg | 16 | 11 | 68.8 | 6 | 37.5 |
| 7 MJ 1999 0.2 mg/Kg + dl propranolol 0.07 mg/Kg | 12 | 10 | 83.3 | 6 | 50 |
| 8 AL 21 011 1.4 mg/Kg | 16 | 12 | 75 | 9 | 56.3 |

The temporal distribution of ventricular ectopic beats following sudden coronary ligation in the present study agrees with that described in anesthetized dogs.¹¹ Multiple factors are probably involved in the production of ectopic impulses after

coronary occlusion. Certainly the catecholamines and the cardiac sympathetic nerves seem to be very important in the genesis of ectopic beats and ventricular fibrillation during the early phases.^{11, 26} Although the doses of beta blocking agents

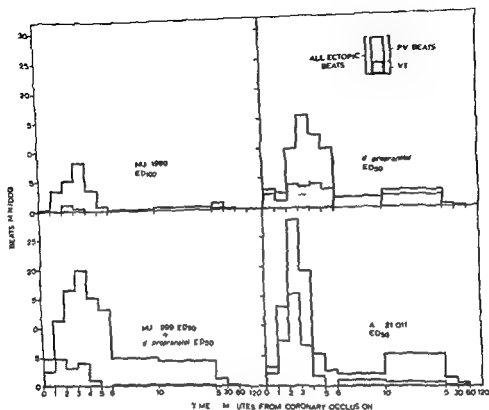


Fig 3B See legend for Fig 3A

Table VI All ventricular ectopic beats (\pm standard error of the mean) per dog per minute following occlusion

| Group | First 5 minutes | | Second 5 minutes | | Third 5 minutes | |
|--|---------------------|----------|--------------------|----------|----------------------|----------|
| | (number/min) | <i>p</i> | (number/min) | <i>p</i> | (number/min) | <i>p</i> |
| 1 Normal saline | 16.06 ± 3.3 | | 4.18 ± 2.13 | | 2.26 ± 2.09 | |
| 2 <i>d</i> propranolol 0.1 mg/kg | 5.12 ± 1.38 | <0.005 | 1.03 ± 0.41 | >0.1 | 0.2 ± 0.1 | >0.3 |
| 3 <i>d</i> propranolol 1.0 mg/kg | 11.32 ± 5.00 | >0.4 | 2.46 ± 1.5 | >0.3 | 1.49 ± 1.01 | >0.7 |
| 4 MJ 1999 0.2 mg/kg | 13.23 ± 4.0 | >0.5 | 2.31 ± 1.27 | >0.5 | 4.84 ± 1.86 | >0.3 |
| 5 MJ 1999 3.2 mg/kg | 4.72 ± 1.59 | <0.005 | 0.16 ± 0.09 | >0.05 | 0.42 ± 0.32 | >0.3 |
| 6 <i>d</i> propranolol 0.07 mg/kg | 6.51 ± 2.77 | <0.05 | 2.99 ± 6.03 | >0.5 | 22.53 ± 21.77 | >0.3 |
| 7 MJ 1999 0.2 mg/kg + <i>d</i> propranolol 0.07 mg/kg | 13.02 ± 4.6 | >0.5 | 3.33 ± 3.8 | >0.5 | 4.24 ± 2.37 | >0.3 |
| 8 A 21 011 1.4 mg/kg | 12.89 ± 3.53 | >0.5 | 1.4 ± 0.82 | >0.5 | 5.33 ± 3.61 | >0.4 |

Table VII Premature ventricular beats (\pm standard error of the mean) per dog per minute

| Group | First 5 minutes | | Second 5 minutes | | Third 5 minutes | |
|--|---------------------|------|--------------------|-------|--------------------|------|
| | (number/min) | p | (number/min) | p | (number/min) | p |
| 1 Normal saline | 5.44 ± 1.44 | | 3.09 ± 1.43 | | 2.26 ± 2.09 | |
| 2 dl propranolol 0.1 mg/kg | 4.26 ± 1.18 | >0.5 | 0.98 ± 0.41 | >0.1 | 0.70 ± 0.10 | >0.3 |
| 3 dl propranolol 1.0 mg/kg | 7.87 ± 2.65 | >0.5 | 2.46 ± 1.51 | >0.7 | 0.51 ± 0.22 | >0.4 |
| 4 MJ 1999 0.2 mg/kg | 5.27 ± 1.52 | >0.9 | 2.06 ± 1.10 | >0.5 | 4.54 ± 1.78 | >0.4 |
| 5 MJ 1999 3.2 mg/kg | 4.47 ± 1.54 | >0.6 | 0.16 ± 0.09 | >0.05 | 0.42 ± 0.32 | >0.3 |
| 6 dl propranolol 0.07 mg/kg | 3.53 ± 1.56 | >0.3 | 5.17 ± 3.26 | >0.5 | 3.20 ± 2.57 | >0.7 |
| 7 MJ 1999 0.2 mg/kg + dl propranolol 0.07 mg/kg | 10.65 ± 3.55 | >0.1 | 6.07 ± 3.82 | >0.4 | 4.04 ± 2.75 | >0.5 |
| 8 AY 21,011 1.4 mg/kg | 6.17 ± 2.00 | >0.7 | 0.68 ± 0.34 | >0.1 | 4.73 ± 3.07 | >0.5 |

selected for use in this study were sufficient to halve or totally inhibit the positive chronotropic response elicited by the injection of 15 μ g per kilogram of body weight isoproterenol they were by no means adequate to block the tachycardia produced immediately following coronary occlusion in the conscious dog. Yet they reduced the incidence of VT significantly. The results thus suggest that the beta blocking capacity of these compounds must involve some process or processes which decreases arrhythmogenicity.²¹ It is well established that dl propranolol has both beta blocking and local anesthetic effects; that its dextroisomer has only local anesthetic action; that MJ 1999 possesses only beta blocking property; and AY 21,011 is cardio selective in its action.²² It is therefore concluded from this study that cardio selective beta blockade alone is sufficient to significantly decrease the incidence of VT following experimental coronary occlusion.

The number of ventricular ectopic beats occurring in trains of four or more (VT) were analyzed separately. The number of ectopic ventricular beats occurring in such trains were significantly reduced ($p < 0.01$) during the first five minutes following coro-

nary occlusion in the dogs given the lower dose of dl propranolol; the higher dose of MJ 1999; the dextroisomer of propranolol alone; and in combination with the lower dose of MJ 1999. It seems possible then that the local anesthetic effect of these drugs in the doses used may have contributed to the reduction of ventricular tachycardia following coronary occlusion. The fact that there were more ectopic beats in the groups treated with the higher dose of dl propranolol is possibly related to the mechanism whereby quinidine like agents produce multifocal ectopic beats.²³

So many patients with coronary artery disease die suddenly (probably in ventricular fibrillation)^{24,25} that the concept of precoronary care²⁶ has evolved. The results of the present study justify further evaluation of the beta adrenergic receptor blocking drugs in experimental coronary occlusion and in carefully selected high risk individuals. An analysis of the present results indicates that the choice of dose may be particularly important and that increasing the dose does not necessarily decrease the occurrence of life threatening arrhythmias.

We wish to thank Dr C. W. Goudey for his continuous encouragement and helpful criticism in

Table VIII Ectopic beats in the form of ventricular tachycardia per dog per minute

| Group | First 5 minutes | | Second 5 minutes | Third 5 minutes |
|--|-----------------|--------|------------------|-----------------|
| | (number/min) | p | (number/min) | (number/min) |
| 1 Normal saline | 10.63 ±2.52 | | 1.07 | 0 |
| 2 dl propranolol 0.1 mg/Kg | 0.86 ±0.76 | <0.001 | 0 | 0 |
| 3 dl propranolol 1.0 mg/kg | 3.45 ±2.83 | >0.05 | 0 | 0.97 |
| 4 MJ 1999 0.2 mg/kg | 8.06 ±3.38 | >0.5 | 2.36 | 0.30 |
| 5 MJ 1999 3.2 mg/kg | 0.25 ±0.19 | <0.001 | 0 | 0 |
| 6 dl propranolol 0.07 mg/kg | 2.98 ±1.74 | <0.05 | 2.81 | 19.33 |
| 7 MJ 1999 0.2 mg/kg + dl propranolol 0.07 mg/kg | 2.36 ±0.97 | <0.001 | 0.27 | 0.2 |
| 8 AL 21 011 1.4 mg/kg | 6.72 ±2.23 | >0.25 | 0.62 | 0.6 |

the preparation of this manuscript and Drs S P Ahuja and M R Roach for their advice. The valuable technical assistance of Mrs V Smith Miss P Stevenson Mr J Klaase and Mr G C Steward is acknowledged. Thanks are also due to Dr R O Davies Ayerst Laboratories Montreal Canada and to Dr G R McKinney Mead Johnson Research Center Evansville Indiana for generous supplies of the blocking agents employed in this study.

REFERENCES

- Somani P and Lum B K B The anti arrhythmic actions of beta adrenergic blocking agents J Pharmacol Exp Ther 147 194 1965
- Pentecost B L and Austen W G Beta adrenergic blockade in experimental myocardial infarction Am Heart J 72 790 1966
- Proger S Sharma A and Naim S Adrenergic beta receptor blockade in experimental myocardial infarction in the dog in The Fifth World Congress of Cardiology New Delhi 1966 National Printing Works Part II p 142
- Shanks H G and Dunlop D Effect of propranolol on arrhythmias following coronary artery occlusion in dogs Cardiovasc Res 11 34 1967
- Dunlop D and Shanks R G Selective blockade of adrenoceptive beta receptors in the heart Br J Pharmacol 32:201 1968
- Kaumann A J and Aramendis P Prevention of ventricular fibrillation induced by coronary ligation J Pharmacol Exp Ther 164 376 1968
- Ceremuzynski L Staszewska Baczak J and Herbaczynska Cedro A Cardiac rhythm disturbances and the release of catecholamines after acute coronary occlusion in dogs Cardiovasc Res 3:190 1969
- Duce B R Garberg I and Smith E R Effects of (plus or minus) propranolol (plus or minus) (plus) and (minus) alprenolol on unanesthetized dogs with ventricular arrhythmias resulting from coronary artery ligation Br J Pharmacol 39 809 1970
- Manning G W McEachern C G and Hall G E Reflex coronary artery spasm following sudden occlusion of other coronary branches Arch Intern Med 61:661 1939
- Bergamaschi M Shanks R G Caravaggi A M and Mandelli V A comparison of the cardiovascular actions of four adrenergic β -receptor blocking agents in resting conscious dogs Am Heart J 82 338 1971
- Harris A S Estandia A and Tillotson W Ventricular ectopic rhythms and ventricular fibrillation following cardiac sympathectomy and coronary occlusion Am J Physiol 162 505 1961
- Khan M I Hamilton J T and Manning G W Protective effect of beta adrenoceptor blockade in experimental coronary occlusion in conscious dogs Am J Cardiol 30 832 1972
- Barrett A M and Cullum V A The biological properties of the optical isomers of propranolol and their effects on cardiac arrhythmias Br J Pharmacol 34 43 1968
- Weiser F Reich L Panagopoulos P Armelini C Gelman M and Kaley G Effects of beta adrenergic blockade by propranolol in experimental acute coronary occlusion with and without hypothermia Fed Proc 25 336 1966
- Jewitt D E Burgess P A and Shillingford

- J P The circulatory effects of Practolol (ICI 50 172) in patients with acute myocardial infarction *Cardiovasc Res* 4:188 1970
- 16 Gibson D and Sowton E Effects of ICI 50172 in man during exercise *Br Med J* 1 213 1968
- 17 Fitzgerald J D and Seales H Effect of a new adrenergic beta blocking agent (ICI 50 172) on heart rate in relation to its blood levels *Int J Clin Pharmacol* 16 467 1968
- 18 Garret J Mahfouz Haptista A and Oswald W Effects of pronethalol on the cardiovascular actions of catecholamines during blockade by phenoxybenzamine *Br J Pharmacol* 27:459 1966
- 19 Fitzgerald J D Perspective in adrenergic beta receptor blockade *Clin Pharmacol Ther* 10 292 1969
- 20 Barrett A M A comparison of the effects of (plus or minus) propranolol and (plus) propranolol in anesthetized dogs: beta receptor blocking and hemodynamic action *J Pharm Pharmacol* 21:241 1969
- 21 Hoffman R P and Grupp G The effects of Sotalol and propranolol on contractile force and atrioventricular conduction time of the dog heart *in situ* *Dis Chest* 55:229 1969
- 22 McLachlan C G Manning G W and Hall G L Sudden occlusion of coronary arteries following removal of cardio-sensory pathways in experimental study *Arch Intern Med* 65 661 1940
- 23 Manning G W and Crudwell G C The effect of demerol ergotamine and dihydroergotamine on mortality after coronary occlusion in dogs *Br Heart J* 9 85 1947
- 24 Clark B B and Cummings J R Arrhythmias following experimental coronary occlusion and their response to drugs *Ann N Y Acad Sci* 64 543 1957
- 25 Jewitt D E Mercer C J Reid D Valori C Thomas M and Shillingford J I Free noradrenaline and adrenaline excretion in relation to the development of cardiac arrhythmias and heart failure in patients with acute myocardial infarction *Lancet* 1 635 1969
- 26 Straszewski Berezak J and Ceremuzynski L The continuous estimation of catecholamine release in the early stages of myocardial infarction in the dog *Clin Sci* 31 531 1968
- 27 Davis L D and Temte J V Effects of propranolol on the transmembrane potentials of ventricular muscle and Purkinje fibers of the dog *Circ Res* 22 661 1968
- 28 Hoffman B F Origin of the heart beat Chap 4 in Luisada A A ed *Cardiovascular functions* Toronto 1967 McGraw Hill Book Co Inc
- 29 Pantridge J F and Geddes J S Mobile intensive-care unit in the management of myocardial infarction *Lancet* 2:271 1967
- 30 Lawrie D M Ventricular fibrillation in acute myocardial infarction *Am Heart J* 8 474 1969
- 31 Lown B and Ruberman W The concept of precoronary care *Mod Concept Cardiovasc Dis* 39 97 1970

Pressure flow studies in man The nature of the aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome*

M Eugene Kendall MD**

Judith C Rembert PhD

Joseph C Greenfield Jr MD***

Durham N C

As early as 1921 Wiggers and Feil¹ described the effect of acute severe mitral insufficiency on aortic blood flow in dogs. Elkins and associates² using an electromagnetic flowmeter to record instantaneous aortic blood flow, noted an abnormally high proportion of flow during the first half of ejection before mitral valve surgery in five patients with severe chronic mitral insufficiency and in nine dogs with acute mitral regurgitation. Except for their work, the nature of blood flow in various types of mitral insufficiency in man has not been evaluated due to the inability to measure phasic aortic blood flow. The pressure gradient technique allows the investigator to estimate the instantaneous aortic blood flow in man.³ Utilizing this technique, the present study was undertaken to evaluate the effects of mitral regurgitation on the pattern of instantaneous aortic blood flow in patients with mitral insufficiency secondary to either rheumatic heart disease, papillary muscle dysfunction, or the prolapsing mitral valve syndrome. Data were ob-

tained during a control state and following pharmacologic interventions with amyl nitrite and phenylephrine.

Methods

Ten male patients and one female patient ranging in age from 41 to 56 years were studied following admission to the Durham Veterans Administration Hospital. Five patients with no demonstrable cardiac disease served as the normal group. In addition, three patients with the classic physical, angiographic, and echocardiographic findings of the prolapsing mitral valve syndrome^{4,5} two patients with severe mitral insufficiency secondary to chronic rheumatic heart disease, and one patient with mitral insufficiency secondary to papillary muscle dysfunction were studied. The clinical data describing the patients with mitral insufficiency are shown in Table 1. Cardiac catheterization was carried out on the normal subjects because of a systolic murmur and they were shown not to have mitral insufficiency or other valvular abnormalities.

From the Department of Medicine (Division of Cardiology), The Veterans Administration Hospital, Durham, N.C. and Duke University Medical Center, Durham, N.C.

Received for publication, October 24, 1972.

Reprints requested to: Joseph C. Greenfield, Jr., MD, Room C-5015, VA Hospital, Durham, N.C. 27705.

*Supported in part by Grant HL-00711 from the National Heart, Lung, and Blood Institute and by Training Grant HL-04807 from the National Institutes of Health.

**Present Address: Cardiology Division, Naval Hospital, Bethesda, Md. 20814.

***Received for publication, December 1, 1972. Address reprint requests to the U.S. Armed Forces Public Health Service.

Table 1 Clinical data

| Patient | Age (yr) | Sex | PA* pressure (mm Hg) | LA pressure (mm Hg) | LVEDP (mm Hg) | Cardiac index (L/min/M ²) |
|---------|----------|-----|----------------------|---------------------|---------------|---------------------------------------|
| R S | 42 | M | 22/8 (14) | (7) | 10 | 3.0 |
| J H | 41 | M | 26/9 (16) | (12) | 16 | 3.3 |
| A W | 41 | F | 36/15 (23) | (12) | 22 | 2.6 |
| M C | 48 | M | 29/12 (18) | — | 8 | 3.0 |
| G G | 56 | M | 37/15 (27) | (26) | 19 | 1.3 |
| O P | 52 | M | 49/24 (40) | (37) | 30 | 2.3 |

Abbreviations and symbols: PA = pulmonary artery; LA = left atrial; LVEDP = left ventricular end diastolic pressure; ECG = electrocardiogram; WNL = within normal limits; LAL = left atrial enlargement; LVL = left ventricular enlargement; () = mean pressure.

*Normal = no evidence of tricuspid valve.

After informed consent was obtained, right and left sided cardiac catheterizations were performed using standard techniques. Left atrial pressures were recorded through a No. 25 Brockenbrough transseptal catheter.* Cardiac outputs were measured using an indicator dilution technique. Left ventricular cineangiography using 75 per cent Renografin† was carried out in all patients to determine the competency of the mitral valve. Mitral insufficiency was graded as mild, moderate or severe. All patients with mitral valve abnormalities had selective coronary arteriograms using the Judkins technique and no lesions were seen except in the patient with papillary muscle dysfunction and posterior wall myocardial infarction. The aortic valve was judged to be normal in all patients. Each patient had an echocardiogram‡ examination using a 2.25 MHz one half inch diameter transducer with a repetition rate of 1,000 per second. The echograms were recorded photographically directly from the oscilloscope.

At the conclusion of the routine catheterization a specially designed 6.5 French Fox-Fry double lumen catheter having lateral pressure tips 4 cm apart was placed percutaneously in the femoral artery and

was advanced to the ascending aorta. The lateral pressures were accurately measured with a transducer amplifier system* and the pressure difference was used to solve the proper equations for phasic blood flow with an analog computer†. Ascending aortic pressure was simultaneously recorded from one lumen of the catheter. Standard Lead II of an electrocardiogram was recorded throughout the study. Continuous phasic aortic flow was then measured in each patient during a control resting state following the inhalation of amyl nitrite and also after the intravenous injection of 0.4 mg phenylephrine. All data were recorded on an optical recorder‡ at a paper speed of 100 mm per second and on FM analog magnetic tape§. The measurements of flow were not associated with any difficulties and the only inconvenience to the patient was the prolongation of the catheterization by approximately 30 minutes.

Phasic blood flow was measured in the ascending aorta by the pressure gradient technique‡. This method is based on the solution of the Navier-Stokes equations of fluid motion which relate the axial

*Model P23Db Statam Instrument Co, Hato Rey, Puerto Rico and Model 350-1500 Hewlett Packard Co, Palo Alto, Calif.

†Model 3400 Donner-Doser System Corp, Concord, Calif.

‡Model 4568B Hewlett Packard Co.

§Model 3955A Hewlett Packard Co.

*Model Catheter and Instrument Corp, Glens Falls, N.Y.

†E.K. Squibb and Sons Inc, N.Y.

‡Echoline 20 Mark II Echogram, Smith-Kline Instruments Inc, Palo Alto, Calif.

| ECG | X ray | Echo | Degree of MI | Functional class (N Y H S) | Diagnosis |
|-----------------------------|---------------------------------|------------------|--------------|----------------------------|--|
| LAD bigeminy | WNL | Prolapsing valve | None | II | Prolapsed mitral valve syndrome |
| WNL | WNL | Prolapsing valve | Moderate | II | Prolapsed mitral valve syndrome |
| WNL | WNL | Prolapsing valve | Severe | III | Prolapsed mitral valve syndrome |
| Posterior infarction | WNL | Normal | Mild | II | Posterior papillary muscle dysfunction |
| Non-specific T wave changes | LAE LVE calcium in mitral valve | Normal | Severe | III | Rheumatic mitral insufficiency |
| LAD LAE LVE | LAE LVE calcium in mitral valve | Normal | Severe | III | Rheumatic mitral insufficiency |

A dog am Ech = echocardiogram III = mitral insufficiency N Y H A = New York Heart Association LAD = left anterior descending

pressure gradient to the flow. The details concerning calibration to obtain flow with this technique have been described elsewhere⁶ and the validity has been demonstrated in a flow generator in the dog aorta and in man.^{1,7,8}

The average values obtained from 10 heart beats in each situation were computed. Stroke volume (cm^3) was obtained by the planimetric integration of the total area under each flow curve. The flow curve was divided into two equal parts as a function of time and the flow during each part was determined and expressed as a per cent of total stroke volume (Fig 1). Peak flow (cm^3 per second) was measured as the maximum value directly from the flow curve. The systolic and diastolic aortic blood pressures (mm Hg) were obtained directly from the record and the mean pressure was computed.

Results

In Table II are listed the hemodynamic parameters for each patient with mitral insufficiency and the mean values for the normal patients. Typical flow curves from one normal and from two patients with the various types of mitral regurgitation in the control state and with each pharmacologic intervention are shown in Fig 1.

The six patients with normal mitral valves had blood flow curves during control and following phenylephrine in which the

per cent flow was approximately evenly divided between the first and second halves of ejection. Inhalation of amyl nitrite resulted in a greater per cent of flow during the first half of ejection.

The proportion of aortic blood flow ejected during the first half of systole in patients with the prolapsing mitral valve syndrome was related to the presence of mitral regurgitation. In the patient (R S) with a prolapsing mitral valve but without mitral insufficiency the blood flow curves were similar to the normal individuals. However in the two patients with mitral regurgitation (J H and A W) the per cent of flow in the first half of ejection was markedly increased during the control state. In these two patients following amyl nitrite and phenylephrine administration the aortic flow during the first half of ejection tended to decrease.

The three patients with either rheumatic mitral insufficiency or papillary muscle dysfunction had a greater proportion of flow during the first half of ejection during control. This augmentation of early systolic flow persisted following amyl nitrite and phenylephrine.

Discussion

The finding of an approximately evenly divided percentage of aortic flow in the first and second halves of ejection in patients with competent mitral valves is in

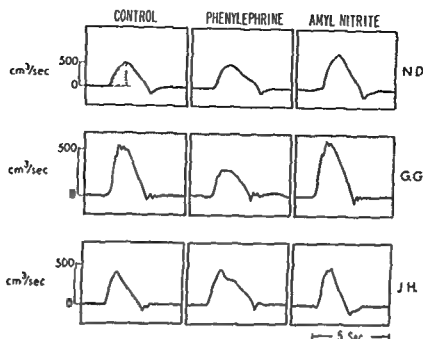


Fig 1 Representative aortic flow curves obtained during the control state following inhalation of amyl nitrite and the intravenous injection of 0.4 mg phenylephrine. *ND* is normal, *GG* has rheumatic mitral insufficiency and *JH* has moderate mitral insufficiency secondary to a prolapsing mitral valve. The stippled area illustrates the flow during the first half of ejection.

keeping with the previous reports from this laboratory^{7,8} and the work of Elkins and associates² following mitral valve replacement. It should be noted that in our normal patients the range of stroke volumes was quite variable, but the per cent of flow in the first half of ejection was consistent from patient to patient during both the control and following administration of amyl nitrite and phenylephrine. Previous data from our laboratory have shown that the proportion of flow during the first half of ejection is not related to the magnitude of the stroke volume.⁷ Inhalation of amyl nitrite by normal individuals decreases mean aortic blood pressure and increases cardiac output by increasing stroke volume with little effect on heart rate.¹⁰ These effects are associated with a greater per cent of aortic blood flow during the first half of ejection (Fig 1). A possible explanation for the phenomenon is that the lower aortic impedance to flow allows a rapid initial ejection resulting in an "empty ventricle" during the latter part of systole. Thus, during the second half of ejection the proportion of flow is reduced. This finding is somewhat analogous to the patients with hypertrophic subaortic stenosis in whom an increase in per cent of flow in the first half of ejection occurs in the control state

possibly due to excessively rapid emptying.⁹ In addition, the decrease in arterial pressure may evoke a release of catecholamines which could result in an increase in myocardial contractility and would augment the ejection during early systole. Phenylephrine usually produces bradycardia, a rise in blood pressure and a decrease in stroke volume. These hemodynamic responses in our normal patients were not marked and the contour of flow curves remained essentially unchanged.

An abnormal pattern of instantaneous aortic blood flow during the control state was demonstrated in each of the patients with mitral insufficiency due to either rheumatic disease or papillary muscle dysfunction in that the per cent of aortic flow is higher during the first half of ejection than in the normal subjects (Fig 1). Similar findings were reported by Elkins and associates² who measured aortic blood flow in five human subjects with an electromagnetic flow meter before and after mitral valve replacement. These workers found 65 per cent of the flow to occur in the first half of ejection prior to the valve replacement and 50 per cent postoperatively. One possible explanation for the finding is that the dilatation of the heart seen in these patients augments the contractile ability of

Table 11 Hemodynamic data

| Patient | | Stroke volume (cm ³) | Peak flow (cm ³ /sec) | Heart rate (beats/min) | Mean aortic blood pressure (mm Hg) | Flow first half ejection (%) |
|---------------|------|-------------------------------------|-------------------------------------|---------------------------|--|------------------------------------|
| R S (I IV) | (C) | 64 | 590 | 75 | 95 | 48 |
| | (I) | 83 | 536 | 62 | 119 | 45 |
| | (AN) | 61 | 580 | 85 | 86 | 55 |
| J H (I IV) | (C) | 60 | 420 | 82 | 109 | 68 |
| | (I) | 75 | 450 | 61 | 120 | 63 |
| | (AN) | 49 | 450 | 109 | 95 | 57 |
| A W (PMV) | (C) | 55 | 390 | 62 | 132 | 66 |
| | (P) | 52 | 300 | 60 | 158 | 59 |
| | (AN) | 62 | 480 | 73 | 101 | 61 |
| M C (PD) | (C) | 52 | 290 | 85 | 104 | 64 |
| | (P) | 65 | 340 | 86 | 114 | 61 |
| | (AN) | 50 | 323 | 100 | 84 | 61 |
| G G (RMI) | (C) | 56 | 500 | 97 | 114 | 58 |
| | (P) | 42 | 360 | 99 | 138 | 56 |
| | (AN) | 60 | 540 | 99 | 103 | 59 |
| O I (RMI) | (C) | 27 | 230 | 97 | 77 | 59 |
| | (P) | 18 | 180 | 92 | 90 | 57 |
| | (AN) | 31 | 250 | 96 | 61 | 57 |
| Normal | (C) | 53 ± 11 | 310 ± 58 | 84 ± 13 | 101 ± 9 | 52 ± 1 |
| | (P)† | 38 ± 11 | 380 ± 57 | 79 ± 14 | 107 ± 9 | 52 ± 2 |
| | (AN) | 76 ± 18 | 490 ± 89 | 86 ± 11 | 73 ± 10 | 60 ± 2 |

†D t = d t h e e s t
Abb t (C) = c t l (P) = p h e s t e p h n e (AN) = m y n i t P M V = p l p g m e a l a l P D = p p l y m u s c l
d y d t R M I = h m i s t e m u s a l u f f i c s

the heart secondary to the Frank Starling mechanism. During the initial part of ejection the flow is increased but as ejection proceeds the volume of the heart decreases markedly due to the large total stroke volume thus during the latter part of ejection the heart cannot maintain this augmented rate of ejection. The observations of Ross and associates¹¹ would tend to substantiate this hypothesis. In these patients the inhalation of amyl nitrite has little effect on the per cent of aortic flow during the first half of ejection because a newly maximal rate of flow for the patient has already been achieved in the control state for the reasons outlined above. The decreased aortic impedance under these circumstances did increase the aortic flow and may have reduced the amount of regurgitation but had little effect on the percentage of flow occurring in the first

half of ejection. Phenylephrine did not change the percentage of aortic flow in the first half of ejection although undoubtedly the total degree of mitral insufficiency was increased.

Late systolic murmurs accompanied by systolic clicks are frequently seen in the prolapsing mitral valve syndrome and are usually associated with mitral insufficiency.^{4, 15, 17} Provocative studies have shown that the late systolic murmur becomes more pansystolic and softer with amyl nitrite and louder without change in systolic timing with pressor amines.¹⁵ The clicks associated with the syndrome often become softer with amyl nitrite and louder with pressor amines. Echocardiography reveals the prolapse of the posterior leaflet of the mitral valve to occur earlier in systole following amyl nitrite.¹⁸ Cineangiographic data reveal earlier and increased mitral

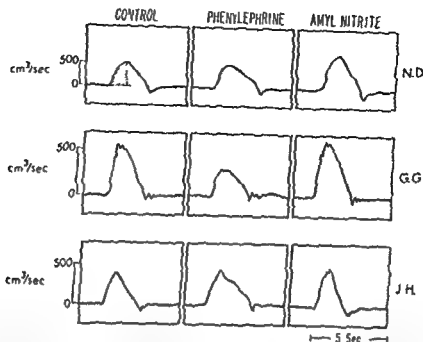


Fig. 1 Representative aortic flow curves obtained during the control state following inhalation of amyl nitrite and the intravenous injection of 0.4 mg. phenylephrine. *N.D.* is normal, *G.G.* has rheumatic mitral insufficiency, and *J.H.* has moderate mitral insufficiency secondary to a prolapsing mitral valve. The stippled area illustrates the flow during the first half of ejection.

keeping with the previous reports from this laboratory^{7,8} and the work of Elkins and associates² following mitral valve replacement. It should be noted that in our normal patients the range of stroke volumes was quite variable but the per cent of flow in the first half of ejection was consistent from patient to patient during both the control and following administration of amyl nitrite and phenylephrine. Previous data from our laboratory have shown that the proportion of flow during the first half of ejection is not related to the magnitude of the stroke volume.⁷ Inhalation of amyl nitrite by normal individuals decreases mean aortic blood pressure and increases cardiac output by increasing stroke volume with little effect on heart rate.¹⁰ These effects are associated with a greater per cent of aortic blood flow during the first half of ejection (Fig. 1). A possible explanation for the phenomenon is that the lower aortic impedance to flow allows a rapid initial ejection resulting in an "empty ventricle" during the latter part of systole. Thus during the second half of ejection the proportion of flow is reduced. This finding is somewhat analogous to the patients with hypertrophic subaortic stenosis in whom an increase in per cent of flow in the first half of ejection occurs in the control state

possibly due to excessively rapid emptying.¹ In addition the decrease in arterial pressure may evoke a release of catecholamines which could result in an increase in myocardial contractility and would augment the ejection during early systole. Phenylephrine usually produces bradycardia, a rise in blood pressure and a decrease in stroke volume. These hemodynamic responses in our normal patients were not marked and the contour of flow curves remained essentially unchanged.

An abnormal pattern of instantaneous aortic blood flow during the control state was demonstrated in each of the patients with mitral insufficiency due to either rheumatic disease or papillary muscle dysfunction in that the per cent of aortic flow is higher during the first half of ejection than in the normal subjects (Fig. 1). Similar findings were reported by Elkins and associates² who measured aortic blood flow in six human subjects with an electromagnetic flow meter before and after mitral valve replacement. These workers found 65 per cent of the flow to occur in the first half of ejection prior to the valve replacement and 50 per cent postoperatively. One possible explanation for the finding is that the dilatation of the heart seen in these patients augments the contractile ability of

- associated with mid systolic click and late systolic murmur *Am J Med* 41:183 1966
- 7 Greenfield J C Jr Harley A Thompson H K and Wallace A C Pressure flow studies in man during atrial fibrillation *J Clin Invest* 47:2411 1968
- 8 Kendall M E Walston A Cobb F R and Greenfield J C Jr Pressure flow studies in man effect of atrial systole on ventricular function in mitral stenosis *J Clin Invest* 50:2053 1971
- 9 Hernandez I R Greenfield J C Jr and McCall B W Pressure flow studies in hypertrophic subaortic stenosis *J Clin Invest* 45:401 1964
- 10 Beck W Schrire V Vogelpoel L Nellen M and Swanepoel A Hemodynamic effects of amyl nitrite and phenylephrine on the normal human circulation and their relation to changes in cardiac murmurs *Am J Cardiol* 8:341 1961
- 11 Ross J Jr Braunwald F and Morrow A G Clinical and hemodynamic observations in pure mitral insufficiency *Am J Cardiol* 2:11 1958
- 12 Levine S and Harvey W P Cardiac murmurs in their clinical auscultation of the heart Philadelphia and London 1959 W B Saunders Co p 194
- 13 Barlow J H Pocock W A Marchand P and Denny M The significance of late systolic murmurs *Am HEART J* 66:443 1963
- 14 Leon D F Leonard J J Kroetz F W Page W L Shaver J A and Lancaster J F Late systolic murmurs clicks and whoops arising from the mitral valve A transseptal intracardiac phonocardiographic analysis *Am HEART J* 72:325 1966
- 15 Ronan J A Perloff J K and Harvey W P Systolic clicks and the late systolic murmur Intracardiac phonocardiographic evidence of their mitral valve origin *Am HEART J* 70:319 1965
- 16 Barlow J B and Bosman C K Aneurysmal protrusion of the posterior leaflet of the mitral valve An auscultatory-electrocardiographic syndrome *Am HEART J* 71:166 1966
- 17 Shell W E Walton J A Clifford M E and Wilks P W III The familial occurrence of the syndrome of mid late systolic click and late systolic murmur *Circulation* 39:327 1969
- 18 Dillon J C Haue C L Chang S and Feigenbaum H Use of echocardiography in patients with prolapsed mitral valve *Circulation* 43:503 1971
- 19 Bittar N and Sosa J A The billowing mitral valve leaflet Report on fourteen patients *Circulation* 38:763 1968

insufficiency secondary to aml nitrite¹⁹

Patient R S had a prolapsing posterior leaflet of the mitral valve as proved by cineangiography and echocardiography but no mitral insufficiency could be seen with angiography. His control aortic flow curves and the responses to both aml nitrite and phenylephrine were similar to the normal patients J H and A W were patients with the prolapsing mitral valve syndrome having moderate and severe mitral insufficiency respectively. During the control state the percentage of flow during the first half of ejection was even more increased in these patients than in the other patients with mitral insufficiency. It is probable that the previously described factors which are responsible for the increase in flow during the first half of ejection in mitral insufficiency also operate in these patients. In addition, since regurgitation across the mitral valve begins later during systole the percentage of aortic flow during the first half of ejection would be further enhanced. An increase in mitral regurgitation during the latter part of systole may also have been present in our patient with papillary muscle dysfunction and may account for the finding of higher proportions of flow during the first half of ejection than in the patients with rheumatic disease. In the two patients with prolapsing mitral valves the inhalation of aml nitrite resulted in a decrease in the per cent of aortic flow during the first half of ejection. Presumably this was due to the occurrence of mitral insufficiency earlier during systole than in the control state as evidenced by the echocardiographic and phonocardiographic findings described above. The reason for the decrease in the per cent of aortic flow during the first half of ejection following phenylephrine administration in these two patients is not immediately apparent. The increased aortic impedance to flow following the drug may result in dilating the left ventricle. This in turn may make the prolapsing valve less insufficient during the latter part of ejection.

Summary

In five normal patients and in six patients with various forms of mitral insufficiency ascending aortic pressure flow relationships

were studied using the pressure gradient technique. Stroke volume, peak flow, mean aortic pressure and the per cent of aortic flow during the first half of ejection were calculated in the control state following aml nitrite inhalation and following the intravenous injection of 0.4 mg of phenylephrine. In normal patients an average of 52 per cent of flow occurred in the first half of ejection both during control and following phenylephrine and increased to 60 per cent after aml nitrite. A patient with the prolapsing mitral valve syndrome but without mitral insufficiency had flow curves similar to the normal. Two patients with moderate and severe mitral insufficiency respectively and a prolapsing mitral valve had a marked increase in flow during the first half of ejection (68 and 66 per cent) during control; the per cent of flow decreased with aml nitrite and phenylephrine administration. Patients with mitral insufficiency secondary to either rheumatic heart disease or papillary muscle dysfunction had 58 to 64 per cent of flow during the first half of ejection during control; this pattern did not change significantly with aml nitrite or phenylephrine.

The author acknowledges the excellent technical assistance provided by Mrs. Kathleen G. Smith, Mrs. Della Foster, Mr. Marvin Micken, Mr. Nobuko U. and Mr. Jesse McCris. The editorial support of Mrs. Hildy Hopkin and Mr. Rosa B. Lathridge and the performing of echocardiogram by Dr. William Spencer. The continued support from Mr. D. Powell and the Medical Illustration Department of the Durham Veterans Administration Hospital is sincerely appreciated.

REFERENCES

1. Wiggers C J and Feil H. Cardiodynamics of mitral insufficiency. *Heart* 9:149, 1921.
2. Elkins R C, Morrow A G, Visko J S and Bruunwald E. The effects of mitral regurgitation on the pattern of instantaneous aortic blood flow. Clinical and experimental observations. *Circulation* 36:45, 1967.
3. Fry D L. The measurement of pulsatile blood flow by the computed pressure gradient technique. *IEEE Trans Bio-Med Eng* 6:259, 1959.
4. Criley J M, Lewis K B, Humphrie J O and Ross R S. Analysis of the mitral valve clinical and cine angiographic findings. *Br Heart J* 28:188, 1966.
5. Greenfield J C Jr. Pressure gradient technique. *Med Res* 11:83, 1966.
6. Hancock F W and Cohen K. The syndrome

- associated with mid systolic click and late systolic murmur *Am J Med* 41:183 1966
- 7 Greenfield J C Jr Harley A Thompson H K and Wallace A G Pressure flow studies in man during atrial fibrillation *J Clin Invest* 47:1411 1968
- 8 Kendall M E Walton A Cobb F R and Greenfield J C Jr Pressure flow studies in man effect of atrial systole on ventricular function in mitral stenosis *J Clin Invest* 50:12613 1971
- 9 Hernandez R R Greenfield J C Jr and McCall B W Pressure flow studies in hypertrophic subaortic stenosis *J Clin Invest* 44:401 1964
- 10 Beck W Schrire V Vogelsoel L Kellen M and Swanepoel A Hemodynamic effects of amyl nitrite and phenylephrine on the normal human circulation and their relation to changes in cardiac murmurs *Am J Cardiol* 8:341 1961
- 11 Ross J Jr Braunwald E and Morrow A G Clinical and hemodynamic observations in pure mitral insufficiency *Am J Cardiol* 2:11 1958
- 12 Levine S and Harvey W P Cardiac murmur in their Clinical auscultation of the heart Philadelphia and London 1959 W B Saunders Co p 194
- 13 Barlow J H Iocock W A Marchand P and Denny M The significance of late systolic murmurs *Am HEART J* 66:443 1963
- 14 Leon D F Leonard J J Kretz F W Page W L Shaver J A and Lancaster J F Late systolic murmurs clicks and whoops arising from the mitral valve A transseptal intracardiac phonocardiographic analysis *Am HEART J* 72:375 1966
- 15 Ronan J A Perloff J K and Harvey W P Systolic clicks and the late systolic murmur Intracardiac phonocardiographic evidence of their mitral valve origin *Am HEART J* 70:319 1965
- 16 Barlow J B and Bosman C K Aneurysmal protrusion of the posterior leaflet of the mitral valve An auscultatory electrocardiographic syndrome *Am HEART J* 71:166 1966
- 17 Shell W E Walton J A Clifford M E and Willis P W III The familial occurrence of the syndrome of mid late systolic click and late systolic murmur *Circulation* 39:1327 1969
- 18 Dillon J C Haine C L Chang M and Feigenbaum H Use of echocardiography in patients with prolapsed mitral valve *Circulation* 43:503 1971
- 19 Bittar N and Sosa J A The billowing mitral valve leaflet Report on fourteen patients *Circulation* 38:763 1968

Anodal stimulation as a cause of pacemaker-induced ventricular fibrillation

Thomas A Preston, MD
Ann Arbor Mich

The possibility of pacemaker induced ventricular arrhythmias has been recognized since the beginning of widespread use of artificial pacemakers^{1,2} but the extent of the clinical risk of such arrhythmias has been questionable.³⁻⁵ The possibility of a dangerous arrhythmia being precipitated by a pacemaker stimulus falling during the vulnerable period of a preceding ventricular beat exists most commonly when an asynchronous artificial pacemaker is in competition with normal sinus rhythm or ectopic beats from any source. The risk is particularly great in patients requiring pacing during acute myocardial infarction as these patients are most susceptible to life threatening ventricular arrhythmias.^{10,20,26,27,28,30,32} This is a review of the reported studies of pacemaker induced ventricular arrhythmias with attention to the type of electrodes used unipolar or bipolar. Every documented case of pacemaker induced ventricular tachycardia or fibrillation occurred in association with a bipolar electrode system, adding support to the concept that these arrhythmias are evoked at the anode and do not occur with unipolar cathodal pacing.

Methods

A survey was made of all reported instances of ventricular tachycardia or fibrillation occurring in human patients with artificial ventricular pacemakers. The events were separated into two categories: (1) documented episodes in which the onset of the ventricular arrhythmia followed a pacemaker stimulus falling during the vulnerable period of the preceding beat, and (2) episodes in which the ventricular arrhythmia but not its onset was documented. There are in addition reports of pacemaker induced ventricular arrhythmias in which the arrhythmia was not documented,^{6,7,12,20,22} or in which the electrode configuration was not stated,^{8,16,17} or the purposes of this study a ventricular arrhythmia was defined as four or more responses following a pacemaker stimulus. The stimulus was judged to fall in the vulnerable period if it occurred during the T wave of the preceding beat.

Each case was further categorized as bipolar or unipolar stimulation with the assumption that in all cases of unipolar pacing, the intracardiac electrode was the cathode and the indifferent electrode was the anode. Although it is possible to re-

From the Department of Medicine, Veterans Administration Hospital, University of Michigan Medical Center, Ann Arbor, Mich.
Received for publication Nov. 6, 1972.
Reprint requests to Thomas A. Preston, MD, Harborview Medical Center, 325 9th Ave., Seattle, Wash. 98104.

Table 1 Documented onset of pacemaker induced ventricular tachycardia or ventricular fibrillation in humans

| | Electrode system | No of patients | Arrhythmia |
|---------------------------|------------------|----------------|------------|
| Bertrand ¹ | Bipolar | 1 | VF |
| Bilitch ²⁰ | Bipolar | 1 | VF |
| Castellanos ¹⁰ | Bipolar | 1 | VF |
| Elmqvist ⁴ | Bipolar | 1 | VF |
| Fletcher ⁴ | Bipolar | 2 | 1 VT 1 VF |
| Grondin ⁷ | Bipolar | 1 | VT |
| Jen ²¹ | Bipolar | 1 | VT |
| Julian ²² | Bipolar | 4 | VF |
| Kleiger ⁷ | Bipolar | 1 | VT |
| Lemberg ²³ | Bipolar | 2 | 1 VT 1 VF |
| Preston ²⁴ | Bipolar | 2 | VF |
| Robinson ⁴ | Bipolar | 1 | VF |
| Roe ¹⁸ | Bipolar | 1 | VF |
| Tavel ¹ | Bipolar | 1 | VF |
| Weinberg ²⁷ | Bipolar | 1 | VT |

Abb 1 VT = ventricular tachycardia VF = ventricular fibrillation

verse polarity in external pacing all commercially available permanent implant pacemakers which are unipolar are fixed with anodal stimulation at the indifferent electrode

Results

In every case of documented onset of ventricular tachycardia or ventricular fibrillation (21 cases) with the pacemaker stimulus falling in the vulnerable period of the preceding beat the electrode system employed was bipolar (see Table 1). We were unable to find such a case using a unipolar system. Sowton and Flores²⁸ have reported two patients who developed ventricular fibrillation shortly after receiving permanent unipolar pacemakers which competed with normal sinus rhythms. In the first patient the onset was not documented although the author considered it to be pacemaker induced. He does state how ever because there is no written ECG record it is just possible that the whole sequence was due to the spontaneous onset of ventricular fibrillation.²⁸ Sowton's second case was also undocumented and although the patient was found with ventricular fibrillation the episode was apparently preceded by abdominal distention vomiting and aspiration. In a patient reported by Roe and Katz²⁹ unipolar pacing did not result in a ventricular arrhythmia whereas

bipolar pacing in the same patient resulted in recurrent bouts of ventricular fibrillation. Of the 21 episodes of pacemaker induced ventricular tachycardia there were no reported episodes of ventricular arrhythmia arising from a pacemaker stimulus falling outside the vulnerable period.

There were 34 episodes of ventricular tachycardia or ventricular fibrillation documented in patients with functioning artificial pacemakers but without documentation of the onset of the arrhythmia.^{1, 2, 4, 10, 11, 14, 17, 18, 21, 22, 23} In 33 patients the pacemaker was operating as a fixed rate (competitive) type and one patient had a normally functioning demand (noncompetitive) unit. Twenty-eight episodes were with bipolar electrodes and six episodes were with unipolar electrodes. Thirty episodes were ventricular fibrillation four episodes were ventricular tachycardia.

Discussion

The problem of pacemaker induced ventricular tachycardia or ventricular fibrillation has been studied since pacemakers have been in common use. Most reports stress the improbability of this phenomenon but some notable exceptions have been documented. Sowton¹ in 1963 reported six patients who developed ventricular tachycardia or ventricular fibrillation while in parasystole with sinus rhythm

and a non inhibited fixed rate pacemaker and concluded that especially in the first two days after implantation there is a risk of pacemaker induced ventricular arrhythmia. Bilitch and co workers²² reported a case of ventricular fibrillation occurring in a patient who had an acute myocardial infarction with documentation of the onset of the arrhythmia at the time of a pacemaker stimulus falling in the vulnerable period of the preceding beat. Castellanos and co workers²⁰ studied 14 human patients and were able to produce repetitive firing in three patients and ventricular fibrillation in one patient by scanning the vulnerable period with bipolar right ventricular pacing. Welti and colleagues²⁰ also produced repetitive firing in humans by scanning the vulnerable period with bipolar stimuli of low amplitude. Lewis and associates⁴¹ using unipolar pacing were unable to produce ventricular arrhythmias.

Clinical studies comparing groups of patients with competitive and noncompetitive systems have yielded conflicting conclusions. Furman and colleagues⁵ report no increased incidence of sudden death in patients with non triggered or competitive units while Bilitch¹⁸ reported death due to ventricular fibrillation in 5 of 40 patients with fixed rate units and only one death due to ventricular fibrillation in 46 patients with demand units. Most investigators feel that pacemaker induced ventricular arrhythmias are most likely during acute myocardial infarction^{19 20 24 26 42} during the first two days after implantation of the electrodes^{3 5} or during periods of hypoxia^{4 27} or electrolyte imbalance²⁷. Another example of ventricular arrhythmia produced by pacing during the vulnerable period is that of paired pacing of the ventricle. Bipolar electrodes are almost always employed for paired pacing and similarly the complication of ventricular tachycardia and fibrillation has been associated with bipolar non unipolar, electrode systems^{40 43 49}.

The results shown in Table I suggest an association of pacemaker induced ventricular fibrillation with bipolar electrodes in the human. There is no documented case of human pacemaker induced ventricular tachycardia or ventricular fibrillation with unipolar electrodes although unipolar systems probably account for almost half of all implanted pacemakers.

Although the cardiac electrode of a permanent unipolar system is connected to the cathode of the impulse generator in temporary systems it is possible to connect the cardiac electrode to the anode of the impulse generator in which case the risk of modulating would be the same as for a bipolar system. This could have been the situation in the two patients reported by Portol and colleagues⁷ and in other cases of temporary unipolar pacing. One of two patients reported by Crandall and co workers²⁴ had a unipolarized electrode system with repetitive firing but whether anode or cathode was made the indifferent electrode is not known.

Some patients with artificial pacing will develop ventricular arrhythmias not induced by a pacing stimulus. For this reason only the documented onset of pacemaker induced ventricular tachycardia or ventricular fibrillation is acceptable as evidence of association with the electrode system in use at the time of the arrhythmia.

Numerous investigators have studied induction of ventricular arrhythmias in animals by repetitive firing during the vulnerable period. Most investigations have been carried out with bipolar electrodes^{51 54 55 61 66 70} with fewer studies of fibrillation thresholds with unipolar cathodal or anodal stimulation^{19 53 59 63 65}. In studies comparing anodal and cathodal stimulation experimental evidence consistently demonstrated increased vulnerability to ventricular arrhythmias from anodal as compared to cathodal stimulation^{42 55 60 64 65}. Chardack and colleagues¹⁹ using unipolar cathodal stimulation were unable to increase the prevalence of ventricular fibrillation in dogs with acute coronary occlusion. Although this was taken as evidence of the safety of artificial pacing it is possible that the low prevalence of fibrillation was associated with the absence of anodal stimulation.

Hoffman and Crane⁴⁴ concluded that during induction of ventricular fibrillation with bipolar electrodes the ventricular fibrillation is initiated at the anode. They state "If the flow of anodal current is sufficiently diffuse it is so difficult to evoke fibrillation that it seems safe to assume that fibrillation evolved by electrical stimulation depends upon anodal".

Cranefield and associates⁵ ruled out a double origin of ventricular fibrillation from both the anode and cathode and concluded that the vulnerable period coincides with the period when threshold excitation occurs at the anode.

Conclusions

The association of bipolar electrodes in humans and animals with pacemaker induced ventricular arrhythmias suggests that in the majority of instances an anode on or within the heart of size and configuration to permit nodal stimulation is necessary to produce ventricular tachycardia or ventricular fibrillation in humans with permanent or temporary pacing systems. As excitation during the vulnerable period is usually possible only at the anode^{6, 7} with the excitation threshold often ten times as great at the cathode it does seem likely that single responses during the vulnerable period are usually evoked at the anode with presently used human pacing systems. It follows that the origin of repetitive ventricular beats or ventricular fibrillation is equally likely to be at the anode.

Evidence is lacking that pacemaker induced arrhythmias are common and commercially available pacemakers have outputs which are probably too small to produce arrhythmias in normal human hearts. It is probable that with impulse durations of one msec or less as is now common with commercial units it is almost impossible to produce arrhythmias due to pacing in the vulnerable period.⁸ However commercial units must be designed to produce stimuli exceeding the excitation thresholds of 95 per cent or more of all patients encountered. With time the excitation threshold for an implanted electrode system rises to 10 times the initial implantation threshold and therefore pacemakers are designed to deliver 10 to 30 times a great stimulus as is required for effective pacing at the time of electrode implantation. Therefore under adverse conditions which lower the fibrillation threshold such as acute myocardial infarction, hypoxia, electrolyte imbalance or during the first two days after implantation pacemaker induced ventricular arrhythmias are possible especially if the anode is on or inside the ventricle.

The use of demand (noncompetitive) pacemakers has reduced the frequency with which pacing stimuli fall in the vulnerable period of the preceding beat but demand type pacemakers are not totally reliable for this purpose as they are subject to failure or outside interference in which case they perform as fixed rate units. Especially with bipolar electrodes even a normally functioning demand pacemaker may fail to sense some spontaneous beats resulting in competitive pacing.

The coronary care unit is a high risk area for pacemaker induced ventricular arrhythmias because of the higher prevalence of ectopic beats which are not sensed by demand pacemakers and the markedly reduced fibrillation threshold of patients with acute myocardial infarction. Therefore in the coronary care unit only unipolar cathodal pacing should be used. As most pacing in a coronary care unit is through temporary catheter electrodes either a remote indifferent electrode (anode) or a catheter with a distal pacing electrode (cathode) and a large proximal electrode (anode—at least ten times the surface area of the distal electrode) positioned outside of the right ventricle should be used. Virtually all temporary endocardial pacing systems employ bipolar electrodes and bipolar pacemakers such that the pacemaker anode can be attached to either electrode. Elimination of bipolar temporary systems should increase the safety of temporary pacing.

Although the danger of pacemaker induced ventricular arrhythmia exists mostly in patients with hypoxia or acute myocardial infarction the greater possibility of such arrhythmias with bipolar electrodes in any situation militates against the general use of bipolar electrodes.

Summary

A review of animal investigations suggests that pacemaker induced ventricular fibrillation usually occurs at the anode and in fact is difficult to evoke at the cathode. A search of the literature showed that every documented episode of pacemaker induced ventricular tachycardia/fibrillation in human has been with a bipolar electrode system. Since the problem most often occurs during temporary pacing associated with myocardial infarction in

and a non inhibited fixed rate pacemaker and concluded that especially in the first two days after implantation there is a risk of pacemaker induced ventricular arrhythmia. Bilitch and co workers²² reported a case of ventricular fibrillation occurring in a patient who had an acute myocardial infarction with documentation of the onset of the arrhythmia at the time of a pacemaker stimulus falling in the vulnerable period of the preceding beat. Castellanos and co workers²⁰ studied 14 human patients and were able to produce repetitive firing in three patients and ventricular fibrillation in one patient by scanning the vulnerable period with bipolar right ventricular pacing. Woltz and colleagues¹⁰ also produced repetitive firing in humans by scanning the vulnerable period with bipolar stimuli of low amplitude. Lewis and associates⁴¹ using unipolar pacing were unable to produce ventricular arrhythmias.

Clinical studies comparing groups of patients with competitive and noncompetitive systems have yielded conflicting conclusions. Furman and colleagues⁵ report no increased incidence of sudden death in patients with non triggered or competitive units while Bilitch¹⁸ reported death due to ventricular fibrillation in 5 of 40 patients with fixed rate units and only one death due to ventricular fibrillation in 46 patients with demand units. Most investigators feel that pacemaker induced ventricular arrhythmias are most likely during acute myocardial infarction^{19, 20, 24, 26, 4} during the first two days after implantation of the electrodes^{2, 23} or during periods of hypoxia^{24, 27} or electrolyte imbalance.⁴² Another example of ventricular arrhythmia produced by pacing during the vulnerable period is that of paired pacing of the ventricle. Bipolar electrodes are almost always employed for paired pacing, and similarly the complication of ventricular tachycardia and fibrillation has been associated with bipolar, not unipolar, electrode systems.^{40, 43, 40}

The results shown in Table I suggest an association of pacemaker induced ventricular fibrillation with bipolar electrodes in the human. There is no documented case of human pacemaker induced ventricular tachycardia or ventricular fibrillation with unipolar electrodes, although unipolar systems probably account for almost half of all implanted pacemakers.

Although the cardiac electrode of a permanent unipolar system is connected to the cathode of the impulse generator in temporary systems it is possible to connect the cardiac electrode to the anode of the impulse generator in which case the risk of nodal pacing would be the same as for a bipolar system. This could have been the situation in the two patients reported by Portal and colleagues⁷ and in other cases of temporary unipolar pacing. One of two patients reported by Grondin and co workers²⁴ had a unipolarized electrode system with repetitive firing but whether anode or cathode was made the indifferent electrode is not known.

Some patients with artificial pacing will develop ventricular arrhythmias not induced by a pacing stimulus. For this reason only the documented onset of pacemaker induced ventricular tachycardia or ventricular fibrillation is acceptable as evidence of association with the electrode system in use at the time of the arrhythmia.

Numerous investigators have studied induction of ventricular arrhythmias in animals by repetitive firing during the vulnerable period. Most investigations have been carried out with bipolar electrodes^{51, 54, 55, 59, 61, 66, 70} with fewer studies of fibrillation thresholds with unipolar cathodal or anodal stimulation.^{19, 53, 63, 62, 63} In studies comparing nodal and cathodal stimulation experimental evidence consistently demonstrated increased vulnerability to ventricular arrhythmias from anodal as compared to cathodal stimulation.^{42, 55, 60, 64, 65} Chardack and colleagues¹⁹ using unipolar cathodal stimulation were unable to increase the prevalence of ventricular fibrillation in dogs with acute coronary occlusion. Although this was taken as evidence of the safety of artificial pacing it is possible that the low prevalence of fibrillation was associated with the absence of anodal stimulation.

Hoffman and Crane⁶⁴ concluded that during induction of ventricular fibrillation with bipolar electrodes the ventricular fibrillation is initiated at the anode. They state: "If the flow of anodal current is sufficiently diffuse it is so difficult to evoke fibrillation that it seems safe to assume that fibrillation evoked by electrical stimulation depends upon anodal..."⁶⁴

- the phenomenon to ventricular fibrillation
Am Heart J 63:367 1962
- 37 Seldon W A and Shanhahn M V Ventricular fibrillation induced by artificial pacemakers
Australas Ann Med 10:92 1966
- 38 Sowton E. Personal communication
- 39 Rose B B and Katy H J Complete heart block with intractable aysiole and recurrent ventricular fibrillation with survival *Am J Cardiol* 15:401 1965
- 40 Welta J Fontaine G and Faquet J Periode vulnerable Risques possibles des rythmes interfeerents et des stimulations paires ou couplees chez l'homme *Arch Mal Coeur* 60:302 1967
- 41 Lewis D H Warner H F and Allan M B Direct measurement of human cardiac excitability (Abstract) *J Clin Invest* 40:1058 1961
- 42 Coumel J, Fobiuso Y Gougon R et al Fibrillation ventriculaire et stimulateurs artificiels *Arch Mal Coeur* 59:919 1966
- 43 Cranefield I F Paired pulse stimulation and postextrasystolic potentiation in the heart *Progr Cardiovasc Dis* 8:346 1966
- 44 Singer D H Golt G and Wagner M L Effects of sustained paired stimulation of the heart in normal dogs and in dogs following coronary artery ligation *Bull N Y Acad Med* 41:652 1965
- 45 Frommer L L Studies on coupled pacing technique and some comments on paired electrical stimulation *Bull N Y Acad Med* 41:670 1965
- 46 Cranefield P F The present status of paired pulse stimulation and postextrasystolic potentiation in the heart *Bull N Y Acad Med* 41:736 1965
- 47 Cranefield I F Scherlag B J Yeh B K et al Treatment of acute cardiac failure by maintained postextrasystolic potentiation *Bull N Y Acad Med* 40:903 1964
- 48 Chardack W M Gage A A and Dean D C Slowing of the heart by paired pulse pacing *Am J Cardiol* 14:374 1964
- 49 Bivwald N S Gay W A Morrow A G et al Sustained paired electrical stimuli *Am J Cardiol* 14:385 1964
- 50 Hsu J, Milozzi A M and Lyon C Ventricular vulnerability to paired pulse stimuli during acute coronary occlusion *Am Heart J* 71:9 1967
- 51 Walker C J The mechanism and nature of ventricular fibrillation *Am Heart J* 70:399 1940
- 52 Walker C J and Wegria R Ventricular fibrillation due to male focalized induction and underer block applied during the vulnerable phase of ventricular systole *Am J Physiol* 128:500 1940
- 53 Walker C J and Wegria R Quantitative measurements of fibrillation thresholds of the mammalian ventricle with observation of the effects of pacing *Am J Physiol* 131:296 1940
- 54 Norrman J C Hys C A Judge R D et al Experimental observation on the instability of totally implantable pacemaker induced ventricular fibrillation *Trans Am Soc Artif Intern Organs* 10:378 1964
- 55 Cranefield I F Hoffman B F and Siebens A V Anodal excitation of cardiac muscle *Am J Physiol* 190:383 1957
- 56 Widmann W D Eisenberg L Levitsky M et al Ventricular fibrillation complicating electrical pacing: Comparison of direct current and radio frequency cardiac pacemaker stimulation *Surg Forum* 14:260 1963
- 57 Hoffman B F Gora E F Wax F S et al Vulnerability to fibrillation and the ventricular excitability curve *Am J Physiol* 167:88 1951
- 58 Chardack W M Heart block treated with an implantable pacemaker *Progr Cardiovasc Dis* 6:307 1964
- 59 Hoffman B F Suckling E F and Brooks C McC Vulnerability of the dog ventricle and effects of defibrillation *Circ Res* 3:147 1955
- 60 Harn A S and Moe G K Idioventricular rhythms and fibrillation induced at the anode or the cathode by direct currents of long duration *Am J Physiol* 136:318 1942
- 61 Moe G K Harn A S and Wiggers C J Analysis of the initiation of fibrillation by electrographic studies *Am J Physiol* 134:473 1941
- 62 Brooks C McC Cranefield P F Hoffman B F et al Anodal effects during the refractory period of cardiac muscle *J Cell Physiol* 48:237 1956
- 63 Van Dam R T Durrer D Strackee J et al Excitability cycle of the dog's left ventricle determined by anodal cathodal and bipolar stimulation *Circ Res* 4:196 1956
- 64 Hoffman B F and Cranefield P F *Electrophysiology of the heart* New York 1960 McGraw Hill Book Company Inc p 225
- 65 Surawicz B Ventricular fibrillation *Am J Cardiol* 28:268 1971
- 66 Barker B M Shine L C Burford T et al Indwelling electronic cardiac pacemakers *JAMA* 186:754 1963
- 67 Rothfeld E L Zucker I R Parsonnet V et al Effect of quinidine on competitive cardiac pacing (Abstract) *Circulation* 32(Suppl 2):182 1965
- 68 Wiggers C J Wegria R and Pinera M The effects of myocardial ischemia on the fibrillation threshold: the mechanism of spontaneous ventricular fibrillation following coronary occlusion *Am J Physiol* 131:309 1940
- 69 Grounau I Lelie G Guignard J et al Evaluation of cardiac drugs in the presence of an electrical pacemaker *J Thorac Cardiovasc Surg* 48:941 1964
- 70 Gerst E Fleming W H and Malon J R Increased susceptibility of the heart to ventricular fibrillation during metabolic acidosis *Circ Res* 19:63 1966
- 71 Bertrand M Cabrol C Guiraudon G et al Fibrillation ventriculaire apres implantation de stimulateur interne *Presse Med* 45:1455 1967
- 72 Ireston T A Unpublished data

polar catheter electrodes should not be used for temporary pacing, and the use of unipolar (cathodal) pacing systems should increase the safety of electrical pacing.

REFERENCES

- Chardick W S, Gage A A and Grentbitch W. Correction of complete heart block by a self contained and subcutaneously implanted pacemaker. *J Thorac Cardiovasc Surg* 42:814 1961
- Bonnabean R C, Bilgutay A M, Sterns I P et al. Observations on sudden death during pacemaker stimulation in complete atrioventricular block. *Trans Am Soc Artif Intern Organs* 9:158 1963
- Sowton L. Artificial pacemaking and sinus rhythm. *Br Heart J* 27:311 1965
- Robinson D E, Lissett H L, Wheeler D H et al. Ventricular fibrillation associated with two functioning implanted cardiac pacemakers. *Am J Cardiol* 15:397 1965
- Tavel M and Lisch C. Repetitive ventricular arrhythmia resulting from artificial internal pacemaker. *Circulation* 30:493 1964
- Bouvrain Y and Slama R. Résultats à long terme de l'entrainement électrocardiologique par stimulateur intracorporel. *Ann Med Interne (Paris)* 116:397 1965
- Forrest R W, Davies J G, Lenthum A et al. Artificial pacing for the heart. *Lancet* 2:1369 1962
- Robinson J S, Sloman G, Hogan J et al. Ventricular tachycardia and fibrillation with implanted electrical pacemakers. *Br Heart J* 27:937 1965
- Noordijk J A, Oey F T J and Tebrin W. Myocardial electrodes and the danger of ventricular fibrillation. *Lancet* 1:975 1961
- Trimble A S. The implantable cardiac pacemaker. Late failures and their management. *J Thorac Cardiovasc Surg* 50:707 1965
- Laurence G H, Laine R M and Hughes M I. Management of complications associated with the use of implantable electronic cardiac pacemakers for the relief of complete heart block. *Am J Surg* 110:177 1965
- Senning A. Problems in the use of pacemakers. *J Cardiovasc Surg* 5:651 1964
- Speir H C, Daugherty D C, Chesney J G et al. An appraisal of the surgical management of heart block. *J Thorac Cardiovasc Surg* 49:743 1965
- Dittmar G A, Friese G and Holder F. Erfahrungen über die langfristige elektrische Reizung des menschlichen Herzens. *Z Kreislaufforsch* 51:66 1967
- Lembert I, Castellanos A and Berkovits B V. Pacemaking on demand in AV block. *JAMA* 191:12 1965
- Elmqvist R, Lundgren J, Lethersson S O, Senning A and Wilhelm Olsson G. Artificial pacemaker for treatment of Adams Stokes syndrome and slow heart rate. *Am Heart J* 65:731 1963
- Sowton L and Flores J. Natural history of pacemaker patients. *Bull N Y Acad Med* 47:999 1971
- Bilitch M. Ventricular fibrillation and pacing. *Ann N Y Acad Sci* 167:934 1969
- Chardick W M, Ishikawa H, Lochler F J et al. Pacing and ventricular fibrillation. *Ann N Y Acad Sci* 167:919 1969
- Julian D G, Valentine P A and Miller G G. Disturbances of rate rhythm and conduction in acute myocardial infarction. *Am J Med* 37:915 1964
- Zoll P. Panel discussion. *Ann N Y Acad Sci* 111:955 1964
- Burchell H B. Analogy of electronic pacemaker and ventricular paroxysm with observations on refractory period, supernormal phase and synchronization. *Circulation* 28:818 1963
- Siddons H and Sowton F. Cardiac pacemakers. Springfield Ill 1967. Charles C Thomas Publisher p 109
- Lurman S and Escher D. Principles and techniques of cardiac pacing. New York 1970. Harper & Row Publishers p 136
- Lurman S, Escher D J and Parker B. The failure of triggered pacemakers. *Am Heart J* 82:28 1971
- Freidberg C K, Cohen H and Donofrio F. Advanced heart block as a complication of acute myocardial infarction. Role of pacemaker therapy. *Progr Cardiovasc Dis* 10:466 1968
- Escher D J W. The present status of clinical cardiac pacing. *Am Heart J* 74:176 1967
- Zucker I R, Rothfeld F I, Furman V, Gilbert L and Bernstein A. Competitive idioventricular and extrinsic pacemaker stimuli in heart block. *Am Heart J* 69:67 1965
- Dixon M E, Trank J W and Dobell R C. Ventricular fibrillation threshold variation with coronary flow and its value in assessing experimental myocardial revascularization. *J Thorac Cardiovasc Surg* 44:670 1964
- MacLenn J D and Phibbs M. Relative effect of chronic ischemia and a myocardial revascularization procedure on the ventricular fibrillation threshold. *Circ Res* 8:473 1960
- Wolff G A, Veith K and Lown B A. A vulnerable period for tachycardia following myocardial infarction. *Cardiovasc Res* 2:111 1968
- Julian D G, Lussers B W and Godman M J. Pacing for heart block in acute myocardial infarction. *Ann N Y Acad Sci* 167:911 1969
- Bilitch M, Cosby R S and Cifferky E A. Ventricular fibrillation and competitive pacing. *N Engl J Med* 276:598 1967
- Dresler W, Jones S and Rubin R. Observations in patients with implanted cardiac pacemakers. Repetitive responses to electrical stimuli. *Am J Cardiol* 15:391 1965
- Freidberg, H D. Syncope during standby cardiac pacing. *Br Heart J* 31:281 1969
- Falmer D G. Interruption of T waves by premature QRS complexes and the relationship of

The value of warning arrhythmias in the prediction of ventricular fibrillation within one hour of coronary occlusion Experimental studies in the baboon*

Koenraad J J Bruynseel MD**

Lionel H Opie MD

Cape Town South Africa

Sudden death in acute myocardial infarction is generally held to be the result of ventricular fibrillation.¹ In acute myocardial infarction the vulnerable period (or descending limb of the T wave) is prolonged and the ventricular fibrillation threshold is decreased both favoring the development of ventricular fibrillation.^{2,3} Primary ventricular fibrillation in acute myocardial infarction has its peak incidence in the first hour and decreases progressively in the next hour and sudden death is the first symptom of ischemic heart disease in 20 to 25 per cent of cases.^{4,5} Of 596 patients with acute myocardial infarction who were not admitted to hospital 75 per cent died within 2 hours of the attack.⁶ A one year survey in the Edinburgh area showed 73 per cent of all deaths occurring within 4 weeks of acute myocardial infarction occurred outside the hospital and 45 per cent of all deaths occurred within 1 hour of the onset of an acute attack.⁷ Thus to investigate the majority of cases of ventricular fibrillation occurring in acute myocardial infarction requires continuous

monitoring of the rhythm in the first hour of the heart attack.

However the average time interval between the onset of illness and admission to the majority of Coronary Care Units is between 6 and 9 hours.^{8,9} Even the fastest mobile Coronary Care Units cannot reach the patient until 20 to 30 minutes have elapsed. These delays explain why many publications on primary ventricular fibrillation in acute myocardial infarction lack information about the first few hours after the onset of the infarction and why there is a controversy about the nature and incidence of warning arrhythmias preceding primary ventricular fibrillation in acute myocardial infarction. Thus the investigation of the incidence of ventricular fibrillation in the first hour of acute myocardial infarction and the identification of warning arrhythmias requires an animal model. We studied a subhuman primate (the Cape Chimera baboon) which we believe provides a better model of the arrhythmias associated with human myocardial infarction than does the dog.¹⁰

From the Ischaemic Heart Disease Laboratory Department of Medicine, Groote Schuur Hospital, University of Cape Town, Cape Town, South Africa.

Received for publication November 6, 1972.

Reprint requests to Dr. L. H. Opie, Groote Schuur Hospital, Cape Town, South Africa.

Part of this report has been presented at the British Cardiac Society Meeting, London, March 1972 and at the Southern African Cardiac Society Meeting, Durban, July 1972.

**Dr. Bruynseel is supported by the Chris Barnard Foundation with contributions from the Cardiac Clinical and Ethical Research Council, the University of Cape Town, and the University of Cape Town.

- 73 Pancredi R G McCallister B D and Mankin H T Temporary transvenous catheter electrode pacing of the heart *Circulation* 36 598 1967
- 74 Grondin J LePage G Karamehmet A et al Pacing induced repetitive firing. Report of two cases *Can Med Assoc J* 96:1477 1967
- 75 Jensen N K Schmidt R Grumelli J J et al Intracavitary cardiac pacing *JAMA* 195 916 1966
- 76 Kleiger R Personal communication
- 77 Weinberg S Personal communication
- 78 Fletcher R Personal communication
- 79 Fulk E A and Hurst J W Complete heart block in acute myocardial infarction *Am J Cardiol* 17 695 1966
- 80 Cristallanos A Lemberg L and Berkovitz B V Repetitive firing during synchronized ventricular stimulation (Abstract) *Am J Cardiol* 17 119 1966

cotomy was performed without coronary artery ligation. In one of these a coronary artery was dissected but was not ligated.

Blood samples The right femoral artery was cannulated and the first blood sample was taken within 30 minutes after the Sernylan injection and before the start of artificial respiration. Thereafter samples were taken throughout the experimental period before and after the production of the infarct.

The initial and final value of pH and P_{CO_2} of all baboons included were within the physiological range and did not differ. The hemoglobin and hematocrit in male baboons were initially 14.7 Cm per 100 ml and 44 per cent and decreased at the end of the experiment by 0.6 Cm per 100 ml and 2 per cent respectively. The hemoglobin and hematocrit in the female baboons were initially 13.8 Cm per 100 ml and 41 per cent and decreased at the end of the experiment by 1.19 Cm per 100 ml and 4 per cent respectively. The greater blood loss in females was due to more prolonged bleeding from the cut sternum. The initial plasma K^+ was 3.9 ± 0.1 mEq per liter which fell by an average of 0.5 mEq per liter during the total period of anesthesia and muscle immobilization which lasted 2 to 3 hours. There was no significant change between the initial and final values of blood enzymes and free fatty acids in all animals. An infusion of 4.5 Cm per liter NaCl was started in 13 baboons 3 minutes after the production of the infarct at an average rate of 0.700 ml per kilogram of body weight per minute until the end of the first hour of observation. In 23 other baboons with infarct no infusion was given during the whole experiment. Of the 10 control baboons 5 received 4.5 Cm per liter NaCl during the first hour and 5 received none. The infusion fluids were preheated to body temperature.

Site of infarction The size of the infarct or the area of the ventricle supplied by the ligated coronary artery was defined by the appearance of a dark brownish blue zone with a sharp edge and marked ST elevation on the epicardial electrocardiogram. At the end of the experiment a transverse cut was made along the clearly defined edge and the infarct was weighed

in the wet state. The size was calculated as the percentage of the total wet heart weight cut 1 cm above the semilunar valves. The infarct area was always transmural and the size was confirmed afterwards by biochemical changes in the tissue (loss of high energy phosphate compounds, glycogen and potassium/sodium ratio) and very low coronary blood flow as measured by injections of microspheres labeled with isotopes.¹¹

Site of infarction The situation of the infarct was anterolateral when the proximal third of the main diagonal branch of the left coronary artery was ligated (Fig. 1). This artery corresponds to the description of the third primary division of the left coronary artery in the baboon.¹² Ligation of the mid third or distal third part of the anterior descending coronary artery resulted in an anteroapical infarct also involving the apex. Ligation of the mid third or distal third part of the circumflex coronary artery produced a posterior infarct. An inferoposterior infarct was obtained by a double ligation of the distal third of the anterior descending and circumflex coronary artery. The right coronary artery and proximal third of the anterior descending and circumflex coronary artery were never tied.

Electrocardiographic monitoring Four needle tipped electrodes of an electrocardiogram recorder (Corbin Farnsworth S&B deluxe scopette with dual trace control with direct graphic recorder ICCR) were connected subcutaneously to the appropriate limbs. A standard lead (III or aVF) was continuously recorded at 5 or 25 mm per second throughout the experiment (1 mV = 1 cm). In the same experiments a tape recorder (Holter System Avionics) was connected by 3 hypodermic needles to the chest for a second continuous recording because the direct writer was intermittently used to record the epicardial electrocardiograph.

Definitions of warning arrhythmias The following arrhythmias were chosen retrospectively after they appeared to be most frequently associated with ventricular fibrillation. R on T phenomenon was defined as a ventricular ectopic beat appearing in the descent of the T wave of a sinus beat or of another ventricular ectopic beat.^{10, 14, 21, 24}

Material and methods

Animals Fifty apparently healthy, wild living Cape Chimpanzee baboons (*Papio Ursinus*) were used for the experiments within 1 or 2 weeks after being captured in their natural surroundings. Forty-two were males with an average body weight of 23.5 ± 0.5 kilograms (mean \pm SEM, range 14 to 32 kilograms) and eight females with an average body weight of 15 ± 1 kilogram (range 9 to 20 kilograms). The ratio of body weight to heart weight was 198 ± 5 ($n = 17$) ± 0.12 gm heart tissue per kilogram of body weight with no significant difference between females and males. The use of the same variety of baboon throughout made it possible to estimate the age by the following criteria:

1. **BODY WEIGHT** Adolescent male and female baboons had a body weight of less than 21 kilograms and 14 kilograms, respectively. Adult and old baboons exceeded these weights.

2. **TEETH** Adolescent male and female baboons had permanent teeth which were small with no signs of wear; no teeth were lacking. The teeth of adult baboons were long, especially the canines. When teeth were lacking, excessively worn, or diseased, these baboons were considered to be old.

3. **BLOOD ALKALINE PHOSPHATASE** The alkaline phosphatase of adolescent baboons was 584 ± 33 IU per milliliter ($n = 9$). Adults and old baboons had an alkaline phosphatase value of 197 ± 23 IU per milliliter ($n = 41$, $p < 0.0005$).

4. **PERICARDIAL FAT AND CORONARY ARTERIES** When the chest was opened, there was generally more yellow epicardial fat in older baboons and their coronary arteries were tortuous. Adolescent baboons were considered to be less than 7 years of age, "adult" baboons between 8 and 20 years of age, and old baboons between 20 to 30 years.¹⁴

Baboons were starved overnight prior to the experiment and received no antibiotic, antituberculous, or antiparasitic treatment nor any other drugs.

Anesthesia and respiration Phencyclidine HCl (Sernylan) 1.25 mg per kilogram of body weight was used to paralyze and to initiate anesthesia which was maintained by intermittent intravenous doses of 30 to 60 mg of pentobarbital (average rate of

administration 0.06 ± 0.003 mg per kilogram of body weight per minute). The baboon was intubated, connected to a Harvard respirator pump (model 607) and ventilated with room air at a fixed rate of 16 strokes per minute against a constant end expiratory pressure of 5 cm water.^{17,18} The stroke volume checked by arterial Astrup readings was about 1 ml per kilogram of body weight. Initial PO_2 was 73 ± 3 mm Hg and the final PO_2 was 82 ± 6 mm Hg (11 baboons).

Operative procedure With the baboon lying supine, the chest was opened by mid-sternal thoracotomy and the heart was suspended in a pericardial cradle after incision of the pericardium parallel to the phrenic nerve. The baboon was turned over to an angle of 45 degrees in the left anterior oblique position for the remainder of the experiment.

The room temperature was kept at about 30° C throughout the whole experiment by electrical fan heating. The body temperature was checked with a thermocouple electrode in the esophagus. The average body temperature was 37° C in males and 35° C in females and decreased by about 0° C by the end of the experimental period.

No antirhythmic drug, analgesic drug, heparin, or oxygen was administered during the experiment.

Production of myocardial infarction One or two branches of the left coronary artery were dissected from their veins and ligated with a linen thread. Within 1 minute of ligation a dark brownish blue epicardial area with a very sharply defined edge appeared. In the next hour of observation the color difference between ischemic and non-ischemic tissue became more marked but no visible change in the size of the infarct occurred. Depending on the size and site of the infarct, bulging of the ischemic free wall of the left ventricle occurred during systole.

Our aim was to produce a single infarct of about 10 per cent of the total wet heart weight. In 34 baboons one ligation of a coronary artery was sufficient but in 6 others 2 contiguous areas supplied by different coronary artery branches had to be tied depending on the anatomy of the coronary arteries. In 10 baboons a thor-

June 86
Volume 3

Table 1 Correlation of size of infarct sex age and mortality rate

| | Male | | Female | |
|--------------------------------------|----------|-----------|----------|----------|
| | Survival | V F† | Survival | V F |
| Adolescent‡ 5-7 year | 2 | 2 | 1 | 0 |
| Adult 8-20 years | 7 | 19 | 3 | 2 |
| Old 20-30 years | 0 | 2 | 0 | 0 |
| Total number | 9 | 23 | 4 | 2 |
| † | 28 | 72 | 66 | 34 |
| Average size infarct (Mean ± SEM) | 7 ± 1°C | 12 ± 15°C | 11 ± 1°C | 12 ± 6°C |

Survival = % of V F at 60 min
V F = ventricular fibrillation at 60 min after coronary ligation
Age group as defined in text
† p < 0.05 for sex difference in comparison
|| p < 0.05

Table 2 Correlation between size and site of infarction and incidence and interval between arterial ligation and development of ventricular fibrillation (V F)

| | Anterolateral infarct | | Anterostapital infarct | | Posterior infarct | | Inferoposterior infarct | |
|--|-----------------------|----------|------------------------|----------|-------------------|----------|-------------------------|----------|
| | V F | Survival | V F | Survival | V F | Survival | V F | Survival |
| Number of baboons | 7 | 0 | 11 | 8 | 4 | 5 | 3 | 1 |
| Size of infarct (Mean ± SEM) | 37 | — | 10 | 11 | 9 | 5 | 12 | 11 |
| Minutes after ligation (Mean ± SEM) | 31 | — | 35 | 60 | 50 | 60 | 23 | 60 |
| † developing V F | 2 | — | 4 | 7 | — | — | 8 | 73 |

Abb. 10 of 11

to the development of ventricular fibrillation in 5 adolescent baboons with infarction 2 baboons developed V F in 31 adult baboons 21 developed V F in 2 old baboons both developed V F (Table 1). Sex was also related to the development of V F. Of 6 female baboons with infarction 3 developed V F. Of 32 male baboons 23 developed V F. These findings correspond well with the correlation of mortality rate with sex and increasing age in humans with acute myocardial infarction^{10,22}. The control baboons consisted of 9 males (2 adolescent 5 adult 1 old with mean body weight of 24.3 ± 1.7 kilograms) and 2

females (1 adolescent weight 12.3 kilograms and 1 adult of 18 kilograms).

The size of the infarct was particularly important for male survival and a 10 per cent infarct was the critical size in this group. The average size of the infarct of the males who developed V F was significantly larger ($p < 0.001$) than of the males who survived.

The female group who survived had a larger infarct than the male survival group ($p < 0.05$). Although some subgroups are rather small it is likely that increasing age and size of infarction increased the incidence of V F for male baboons. Females

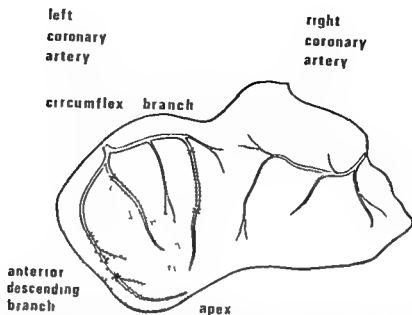


Fig. 1 Schematic representation of the free wall of the left and right ventricle spread open. The shaded areas on the left side correspond to the infarction areas produced by ligation of the mid and distal third portions of the descending coronary branch. The shaded areas on the right side correspond to the infarction areas produced by ligation of the mid and distal third portions of the circumflex coronary branch. The shaded area in the left ventricle corresponds to the infarction area produced by ligation of the proximal third of the main diagonal branch (third primary branch).

R on T beats initiating ventricular tachycardia or ventricular fibrillation were not included as a warning arrhythmia. Ventricular bigeminy was defined as the alternation of a sinus beat and a premature ventricular ectopic beat for at least 4 consecutive beats.¹⁴ Ventricular tachycardia was defined as 3 or more unifocal or multifocal consecutive ventricular ectopic beats at a rate of more than 140 per minute.^{2,14,25,6} Short bursts as well as long runs were included. Ventricular tachycardia which initiated ventricular fibrillation was not included. Occasional arrhythmias occurring during the intermittent recording of the electrogram with the handheld epicardial electrode were discarded.

The method of prediction of imminent ventricular fibrillation in the presence of acute myocardial infarction was based on the first and last appearance time of the above three arrhythmias. The first appearance time was defined as the time interval between the beginning of the first occurrence of the arrhythmia after coronary artery ligation and the end of the experiment. The last appearance time was defined as the time interval between the beginning of the last occurrence of the arrhythmia (before the end of the experiment) and the

end which was onset of ventricular fibrillation or the 60th minute after coronary artery ligation.

Discarded experiments Two baboons in whom accidental ventricular fibrillation was induced mechanically by the application of the epicardial electrode to the boundary of the infarct were discarded from the results.

Expression of data Calculations of group differences paired data analysis and their statistical significance were performed by standard statistical methods.²⁷ Results are expressed as means \pm SEM (number of observations).

Results and discussion

Of the 38 baboons with acute myocardial infarction 21 (66 per cent) spontaneously developed ventricular fibrillation (VF). On all occasions VF did not spontaneously revert and was not artificially interrupted. All 10 control baboons survived 1 hour of observation. The results are analyzed in relation to the incidence of VF or survival for the first hour after coronary artery ligation or in the case of sham operations for a control period of one hour.

AGE AND SEX OF ANIMALS AND SIZE AND SITE OF INFARCTION Age was related

V 1 me 86
N mde 3

Table V Relation between ventricular bigeminy (VB) and incidence of ventricular fibrillation (VF)

| | Group with VF† | Group with 1 hr survival | p value |
|--|----------------|--------------------------|-------------|
| Number of baboons | 17 | 6 | |
| Total number of episodes of VB | 36 | 10 | |
| Incidence | 36/17 = 2.11 | 10/6 = 1.66 | |
| Coupling interval per animal | | | |
| Variable | 5 | 0 | |
| Constant | 12 | 6 | |
| First time of appearance of VB before end in min | 16 | 34 | |
| Mean \pm SEM | ± 2 | ± 5 | $p < 0.001$ |
| Number | 17 | 6 | |
| Last time of appearance of VB before end in min | 5 | 19 | |
| Mean \pm SEM | ± 2 | ± 6 | $p < 0.001$ |
| Number | 17 | 6 | |

VB = ventricular bigeminy

VF = ventricular fibrillation

fibrillation (Table IV) but this is probably due to age since we found no spontaneously hypertrophied or dilated hearts in our series of healthy baboons. It is known that a certain critical mass of myocardium is necessary to maintain the fibrillation activity in the ventricle.^{22,23}

2 HEART RATE Before thoracotomy, the initial heart rate was 95 ± 4 beats per minute for the over all group which corresponds well with the telemetrically measured rate of 82 beats per minute in 4 awake adult baboons who were at rest.²⁴ The heart rate after thoracotomy and before coronary artery ligation was 102 ± 4 beats per minute. The increase of heart rate was due to the surgical procedure with further increases of heart rate due to intravenous injections of pentobarbital given to maintain anesthesia. The final heart rate of 121 ± 4 beats per minute in surviving baboons corresponded to the rate in mildly exercised baboons.²⁵ There was no difference between the heart rate of the baboons of the control group (no arterial ligation) and the baboons who developed ventricular fibrillation or survived the first hour and VF occurred at any heart rate between 80 and 200 sinus beats per minute.

3 BRADYARRHYTHMIAS The rarity of bradyarrhythmias and the absence of asystole corresponded with the findings of other investigators^{21,27} and is probably ex-

plained by the fact that we never ligated the right coronary artery and its branches to the sinus node and the atrioventricular node. Nor did we ligate the initial part of the anterior descending coronary artery with its multiple perforating ventricular septal branches nor the proximal third of the circumflex coronary artery with its left atrial branch. Bradyarrhythmia did not precede primary ventricular fibrillation which is in agreement with the findings of some authors^{28,29} and in contradiction to the findings of others.^{14,30,31}

4 ARRHYTHMIAS WARNING OF VENTRICULAR FIBRILLATION Ventricular bigeminy, ventricular tachycardia and R-on-T beats were the three most common arrhythmias preceding ventricular fibrillation and frequently appeared on several occasions in the same animal. The duration of the attack of ventricular bigeminy was the longest in the group which developed VF (Table V).

The incidence or the number of ventricular bigeminy attacks per animal was not helpful in distinguishing survival from VF. There was however a significant difference between the appearance time of the attack of ventricular bigeminy comparing the group who survived and the group who developed VF ($p < 0.001$). Ventricular bigeminy with a variable coupling interval was always followed by VF. Ventricular

Table III Correlation between number of ligations size of infarct, and incidence of survival

| Number of ligations | Group with VF* | | Group with survival for 1 hour | | % surviving for 1 hour |
|---------------------|----------------|-------------------------------|--------------------------------|-------------------------------|------------------------|
| | Number | Infarct size (Mean \pm SEM) | Number | Infarct size (Mean \pm SEM) | |
| 0 | 0 | — | 10 | — | 100 |
| 1 | 19 | 12 \pm 1% ₀ | 12 | 8 \pm 1% ₀ | 44 |
| 2 | 6 | 11 \pm 2% ₀ | 1 | 11% ₀ | 14 |

*Abbreviations VF = ventricular fibrillation

Table IV Relation between size of infarct total wet weight of heart and mortality rate

| | Control group | Infarct group with survival for 1 hour | Infarct group with VF* |
|--|---------------|--|------------------------|
| Total wet weight of heart in grams | | | |
| (Mean \pm S.E.M) | 114 \pm 14 | 99 \pm 10 | 123 \pm 7 |
| Number | (10) | (13) | (25) |
| P value | | p > 0.05 | p < 0.05 |
| Size of infarct as % of total heart weight | — | 8.3 \pm 0.9 (13) | 11.9 \pm 1.1 (25) |
| | | | p < 0.01 |

*VF = ventricular fibrillation

survived larger infarcts than males. Other authors¹⁰ found a good correlation between the size of the infarct and the incidence of VF in dogs and pigs.

In the group with *anterolateral* infarct all 7 baboons developed VF. The average size of the infarct was significantly larger than the overall size of the infarct of all baboons with VF ($p < 0.025$) (Table II).

The average size of the infarct in the group with *inferior* infarct was identical in the groups who survived or developed VF; thus parameters other than size and site of infarct must have operated. The group of baboons with a *posterior* infarct and VF had the same size of infarct (nearly 10 per cent) as most other groups but developed VF later than all other groups with infarct and VF ($p < 0.05$).

In the group with *inferoposterior* in

farct 3 of 4 of the baboons developed VF. We believe that anterolaterally situated infarcts result in large sized infarcts with increasing risk of developing VF.¹¹ The largest sized infarcts with survival were found in the anteroseptal apical area. These findings contrast with the findings in the greyhound dogs in whom *anterolateral* infarcts were relatively free from the development of ventricular fibrillation whereas fibrillation occurred more readily in *anteroseptal* infarctions.¹²

The average time of onset of VF was 30 to 35 minutes after the ligation of the coronary artery. Infarcts due to *double occlusion* of two coronary artery branches with contiguous blood supply had only a 14 per cent survival rate for an average sized infarct (Tables II and III).

The total wet heart weight was directly proportional to the incidence of ventricular

Table V Relation between ventricular bigeminy (V B) and incidence of ventricular fibrillation (V F)

| | Group with V F† | Group with 1 hr survival | p value |
|---|-----------------|--------------------------|-----------|
| Number of baboons | 17 | 6 | |
| Total number of episodes of V B | 36 | 10 | |
| Incidence | 36/17 = 2.11 | 10/6 = 1.66 | |
| Coupling interval per animal | | | |
| Variable | 5 | 0 | |
| Constant | 12 | 6 | |
| First time of appearance of V B before end in min | III | 31 | |
| Mean = SEM | ± 2 | ± 5 | p < 0.001 |
| Number | 17 | 6 | |
| Last time of appearance of V B before end in min | 5 | 29 | |
| Mean = SEM | ± 2 | ± 6 | p < 0.001 |
| Number | 17 | 6 | |

*V B = ventricular bigeminy
†V F = ventricular fibrillation

fibrillation (Table IV) but this is probably due to age since we found no spontaneously hypertrophied or dilated hearts in our series of healthy baboons. It is known that a certain critical mass of myocardium is necessary to maintain the fibrillation activity in the ventricle.^{24,25}

2. HEART RATE Before thoracotomy the initial heart rate was 90 ± 4 beats per minute for the over all group which corresponds well with the telemetrically measured rate of 82 beats per minute in 4 awake adult baboons who were at rest.²⁵ The heart rate after thoracotomy and before coronary artery ligation was 102 ± 4 beats per minute. The increase of heart rate was due to the surgical procedure with further increases of heart rate due to intravenous injections of pentobarbital given to maintain anesthesia. The final heart rate of 121 ± 4 beats per minute in surviving baboons corresponded to the rate in mildly exercised baboons.²⁵ There was no difference between the heart rate of the baboons of the control group (no arterial ligation) and the baboons who developed ventricular fibrillation or survived the first hour and V F occurred at any heart rate between 80 and 200 sinus beats per minute.

3. BRADYARRHYTHMIAS The rarity of bradyarrhythmias and the absence of asystole corresponded with the findings of other investigators^{26,27} and is probably ex-

plained by the fact that we never ligated the right coronary artery and its branches to the sinus node and the atrioventricular node. Nor did we ligate the initial part of the anterior descending coronary artery with its multiple perforating ventricular septal branches nor the proximal third of the circumflex coronary artery with its left atrial branch. Bradyarrhythmia did not precede primary ventricular fibrillation which is in agreement with the findings of some authors^{28,29} and in contradiction to the findings of others.^{30,31,32}

4. ARRHYTHMIAS WARNING OF VENTRICULAR FIBRILLATION Ventricular bigeminy, ventricular tachycardia and R-on T beats were the three most common arrhythmias preceding ventricular fibrillation and frequently appeared on several occasions in the same animal. The duration of the attack of ventricular bigeminy was the longest in the group which developed V F (Table V).

The incidence or the number of ventricular bigeminy attacks per animal was not helpful in distinguishing survival from V F. There was however a significant difference between the appearance time of the attack of ventricular bigeminy comparing the group who survived and the group who developed V F ($p < 0.001$). Ventricular bigeminy with a variable coupling interval was always followed by V F. Ventricular

Table III Correlation between number of ligations, size of infarct, and incidence of survival

| Number of ligations | Group with VF* | | Group with survival for 1 hour | | % surviving for 1 hour |
|---------------------|----------------|-------------------------------|--------------------------------|-------------------------------|------------------------|
| | Number | Infarct size (Mean \pm SEM) | Number | Infarct size (Mean \pm SEM) | |
| 0 | 0 | — | 10 | — | 100 |
| 1 | 19 | 12 \pm 1% | 12 | 8 \pm 1% | 44 |
| 2 | 8 | 11 \pm 2% | 1 | 11% | 14 |

*Abbreviations: VF = ventricular fibrillation

Table IV Relation between size of infarct, total wet weight of heart, and mortality rate

| | Control group | Infarct group with survival for 1 hour | Infarct group with VF* |
|--|---------------|--|------------------------|
| Total wet weight of heart in grams | | | |
| (Mean \pm SEM) | 114 \pm 14 | 99 \pm 10 | 123 \pm 7 |
| Number | (10) | (13) | (75) |
| P value | | $p < 0.05$ | |
| Size of infarct as % of total heart weight | — | 8.3 \pm 0.9 (13) | 11.9 \pm 1.1 (25) |
| | | $p < 0.01$ | |

*VF = ventricular fibrillation

survived larger infarcts than males. Other authors¹⁰ found a good correlation between the size of the infarct and the incidence of VI in dogs and pigs.

In the group with anterolateral infarct all 7 baboons developed VI. The average size of the infarct was significantly larger than the overall size of the infarct of all baboons with VI ($p < 0.025$) (Table II).

The average size of the infarct in the group with inferior infarct was identical in the groups who survived or developed VF, thus parameters other than size and site of infarct must have operated. The group of baboons with a posterolateral infarct and VF had the same size of infarct (nearly 10 per cent) as most other groups but developed VF later than all other groups with infarct and VF ($p < 0.05$).

In the group with inferoposterior in-

farct 3 of 4 of the baboons developed VF. We believe that anterolaterally situated infarcts result in large sized infarcts with increasing risk of developing VI.¹¹ The largest sized infarcts with survival were found in the interoseptal apical area. These findings contrast with the findings in the greyhound dogs in whom anterolateral infarcts were relatively free from the development of ventricular fibrillation whereas fibrillation occurred more readily in anteroseptal infarctions.⁴

The average time of onset of VI was 30 to 35 minutes after the ligation of the coronary artery. Infarcts due to double occlusion of two coronary artery branches with contiguous blood supply had only a 14 per cent survival rate for an average sized infarct (Tables II and III).

The total wet heart weight was directly proportional to the incidence of ventricular

Table VIII Correlation of warning arrhythmias with incidence of ventricular fibrillation and survival rate

| | Group with 1 F | | Group with 1 hr survival | |
|-----------------------|----------------|---|--------------------------|----|
| Warning arrhythmia | 24 | a | b | 6 |
| No warning arrhythmia | 1 | c | d | 17 |

$$\frac{[(d-b) - 1/2]^2}{(a+b)(c+d)(1+)(1+d)} = 22.2 \quad p < 0.001 \quad (n=48)$$

$$\text{Coefficient of association } \frac{d-b}{d+b} = 0.975$$

$$\text{Coefficient of relative risk } \frac{d-b}{\sqrt{(a+b)(c+d)(1+)(1+d)}} = 0.725$$

Table IX Frequency of ventricular fibrillation following each warning arrhythmia

| Warning arrhythmia | Number of baboons with VF* | Number baboons with survival for 1 hour | % developing 1 F |
|---|----------------------------|---|------------------|
| R on T | 9 | 4 | 69 |
| Ventricular bigeminy | 17 | 6 | 74 |
| Ventricular tachycardia | 11 | 1 | 91 |
| More than 5 premature ventricular ectopic beats/min | 7 | 2 | 77 |

*VF = ventricular fibrillation.

Table X Relation between number of warning arrhythmias and incidence of ventricular fibrillation

| Number of arrhythmias | Total number of baboons | Number with VF† | % 1 F |
|-----------------------|-------------------------|-----------------|-------|
| 0 | 18 | 1 | 5 |
| 1 | 17 | 15 | 88 |
| 2 | 8 | 5 | 63 |
| 3 | 5 | 4 | 80 |

*% of baboons with ventricular tachycardia or R-T phenomenon.
†VF = ventricular fibrillation.

An approximate quantitative assessment of the ventricular irritability is derived from the average time interval between the first time of the appearance of each arrhythmia and the time of the onset of ventricular fibrillation. This time interval is the prediction time for imminent ventricular fibrillation for each arrhythmia and is obtained when the average time interval between the first appearance of each

arrhythmia and the time of onset of VF is taken and twice the standard time error is added. The practical value of this observation is that ventricular tachycardia, ventricular bigeminy, and R on T beats in the presence of acute myocardial infarction lose their value in predicting imminent VF once more than 30, 20, or 10 minutes (respectively) have elapsed after the arrhythmia has commenced.

Table VI Relation between ventricular tachycardia (VT) and incidence of ventricular fibrillation (VF)

| | Group with VF* | Group with 1 hr survival |
|---|----------------|--------------------------|
| Number of baboons | 11 | 1 |
| Total number of episodes of VT† | 47 | 3 |
| Duration of episodes of VT in sec (Mean \pm SEM) | 5 \pm 2 | 15 \pm 0.3 |
| First time of appearance of VT before end in min (Mean \pm SEM) | 21 \pm 5 | 50 |
| Number | (11) | (1) |
| Last time of appearance of VT before end in min (Mean \pm SEM) | 6 \pm 2 | 50 |
| Number | (11) | (1) |

*VF = ventricular fibrillation

†VT = ventricular tachycardia

Table VII Relation between R on T phenomenon and incidence of ventricular fibrillation (VF)

| | Group with VF* | Group with 1 hr survival | P value |
|---|--------------------|--------------------------|------------|
| Number of baboons | 9 | 4 | |
| Total number of R on T beats | 15 | 4 | |
| First time of appearance of R on T before end in min (Mean \pm SEM) | 4.5 \pm 3 | 36 \pm 7 | p < 0.0075 |
| Number | (9) | (4) | |
| Last time of appearance of R on T before end (Mean \pm SEM) | 67 sec \pm 15 | 36 min \pm 7 | p < 0.0075 |
| Number | (9) | (4) | |

*VF = ventricular fibrillation

tachycardia was highly specific in predicting VF (Table VI)

The development of R on T beats could be followed by VF within 1 to 5 min thereafter VF did not occur frequently ($p < 0.0025$, Table VII). The presence of one or more of these 3 warning arrhythmias or the absence of all 3, correctly predicted imminent VF or survival for 1 hour in 41 of 48 baboons (85 per cent) after myocardial infarction or sham operation (Table VIII). It was possible to predict VF in 24 of the 25 who developed VF—the accuracy was 96 per cent. Survival after 1 hour was correctly predicted in 17 of 23 baboons (74 per cent accuracy). The inci-

dence of the 3 warning arrhythmias thus somewhat overestimated the risk of imminent VF. All 10 control baboons survived; one of these developed ventricular bigeminy which lasted for 2 seconds.

Each warning arrhythmia itself had about a 75 per cent incidence of VF, and the appearance of more than one arrhythmia per animal did not increase the risk of developing VF (Tables IX and X). The appearance of these 3 arrhythmias in the presence of acute myocardial infarction indicates electrical disturbances of the ischemic ventricle and all have a similar qualitative value in estimating developing electrical instability.

in predicting the time that ventricular fibrillation subsequently develops

Summary

Factors associated with the development of ventricular fibrillation after coronary artery ligation were studied in a subhuman primate (Cape Chacma baboon). In 25 or 66 per cent of 38 baboons primary ventricular fibrillation occurred within the first hour after the onset of acute myocardial infarction. Increasing age, total heart weight and the size of the infarct were directly related to the incidence of primary ventricular fibrillation. Antero-lateral infarcts had the highest risk of ventricular fibrillation. Anteroseptal and posterior infarcts had the best survival rate for the first hour. Male baboons were more prone to develop ventricular fibrillation than were females. There was no definite progression from ventricular ectopic beats to ventricular fibrillation. In the presence of ventricular tachycardia (even when brief in duration), ventricular bigeminy or R on T beats, ventricular fibrillation has to be expected from the time of onset of the arrhythmia till 30, 20 or 10 minutes have elapsed respectively. Beyond these times ventricular fibrillation did not develop during the experimental period. Conversely the absence of these signs could predict survival for 1 hour. The overall efficiency of the warning signs in predicting ventricular fibrillation or survival was 85 per cent. Ventricular fibrillation occurred without any of these 3 warning signs in only 1 baboon (5 per cent of all cases). It is suggested that these warning arrhythmias could have a practical value in the management of patients with acute myocardial infarction of recent onset by anticipating the time of impending ventricular fibrillation.

Prof. Dr J. E. Kench and Dr E. Symson are thanked for blood analyses. We also thank Mr Petrus Kew and Mr M. Parker for technical assistance and Dr W. Lubbe for collaborative work.

REFERENCES

1. Lowy B and Wolf M. Approaches to sudden death from coronary heart disease. *Circulation* 44:130 1971.
2. Han J. Ventricular fibrillability during acute

- coronary occlusion. *Am J Cardiol* 21:857 1969.
3. Wiggers C J, Wegria R and Linera B. The effects of myocardial ischaemia on the fibrillation threshold. The mechanism of spontaneous ventricular fibrillation following coronary occlusion. *Am J Physiol* 131:1309 1940.
4. Lovel R H and Liners R J. Mechanisms of sudden death and their implications for prevention and management. *Proc Cardiovasc Dis* 13:182 1971.
5. Gordon T and Kannel W B. Premature mortality from coronary heart disease. *JAMA* 218:1617 1971.
6. McNeill R H and Lembergen J. Duration of last attack in 998 fatal cases of coronary artery disease and its relation to possible cardiac resuscitation. *Br Med J* 3:139 1968.
7. Armstrong A, Duncan H, Oliver M F, Julian D G, Donald K W, Fulton M, Lutz W and Morrison V L. Natural history of acute coronary heart attacks. *Br Heart J* 34:67 1972.
8. Goldstein S, Greene W and Moss A J. Sudden death and hospitalization delay in acute myocardial infarction. *Am J Cardiol* 27 (Abstract) 266 1972.
9. Pentecost B L and Mayne V M C. Results of a general hospital coronary care unit. *Br Med J* 1:1830 1968.
10. Lawrie D M, Greenwood T W, Goddard M, Harvey A C, Donald K W, Julian D G and Oliver M F. A coronary care unit in the routine management of acute myocardial infarction. *Lancet* 2:109 1967.
11. Darby S, Bennett M A, Cruickshank J C and Pentecost B L. Trial of combined 1 M and 1 V lignocaine in prophylaxis of ventricular tachyarrhythmias. *Lancet* 1:817 1972.
12. Church G and Biern R O. Intensive coronary care: a practical system for a small hospital without house staff. *N Engl J Med* 281:1155 1969.
13. Wyman M G. Prevention of primary ventricular fibrillation in acute myocardial infarction. *Am J Cardiol* 29 (Abstract) 298 1972.
14. Dhurandhar R W, McMillan R L and Brown K W G. Primary ventricular fibrillation complicating acute myocardial infarction. *Am J Cardiol* 27:347 1971.
15. Bruynel K and Opie L H. The baboon as an experimental animal for the production of myocardial infarction. Comparison with the mongrel dog. In Oliver M F and Julian D G (editors). Effect of acute ischaemia on myocardial function. Edinburgh 1972. E & S Livingstone Ltd. (In press).
16. Newsome J. Baboons. The UFAW handbook on the care and management of laboratory animals. Edinburgh 1967. E & S Livingstone Ltd. p 709.
17. Woods R, Roberts R M, Sugarman R J and Tour L M K R. Our inexpensive continuous positive and expiratory pressure (PEEP) adaptor for positive pressure respirators. *Chest* 61:316 1972.

A frequency of more than 5 isolated premature ventricular ectopic beats per minute, appearing after the descent of the T wave, is a well known sign indicating electrical instability after acute myocardial infarction. However in our series this sign was less frequent than ventricular bigeminy and tachycardia or the R on T phenomenon (Table IX). When it occurred it was always associated with at least one of the 3 previously mentioned arrhythmias and adding the frequency of ectopic beats did not alter the efficiency of prediction of imminent V F. Bennett and Pentecost,⁴² in a recent study in patients in a Coronary Care Unit, and Moss and associates³⁹ did not find the frequency of single ventricular ectopic beats to be a good warning sign. Furthermore single ventricular ectopic beats may only have a warning value if they originate in the left ventricle.^{14,43} In practice it is difficult to site ventricular origin of an isolated ectopic beat with the single classic precordial bipolar monitor lead used in coronary care units: generally two simultaneously recorded standard leads are necessary to place the origin of the ventricular ectopic focus. One minute of continuous electrocardiographic recording or counting is often necessary to detect the arrhythmia. This procedure is more time consuming than detecting the other 3 warning arrhythmias selected here.

Other arrhythmias such as sinus tachycardia, ventricular trigeminy, and bucket rhythm⁴⁴ were infrequently present. No constant pattern of rhythmicity of ventricular arrhythmias or progressive increase of the quantity of ventricular ectopics preceding ventricular fibrillation could be detected.

Conclusion

Previous authors have found that the period of susceptibility to early ventricular fibrillation in dogs with acute coronary artery occlusion was short in duration and that fibrillation was initiated by a paroxysm of ventricular ectopic beats accelerating in frequency.^{45,46} These authors drew attention to the causal relationship between progressive increase of ventricular premature beats and the initiation of ventricular fibrillation. However, this pattern

of events was not usual in our study of the baboon, and is also infrequent in patients with acute myocardial infarction.^{11,42} Thus a search had to be made for other warning signs of ventricular fibrillation.

Hill and colleagues²⁶ found 57 episodes of primary ventricular fibrillation in 18 of 26 rhesus monkeys within the first 40 minutes of the onset of acute myocardial infarction. Each time ventricular fibrillation was electrically stopped. Apart from premature ventricular ectopic beats immediately after the coronary artery occlusion, they found no other complex ventricular arrhythmias preceding V F. We believe that this apparent lack of warning signs is due to the marked increase of sinus rhythm (200 to 240 beats per minute) of the conscious animals suppressing any slower ventricular focus and to the fact that no continuous ECG rhythm recording was undertaken during the first hours after the acute myocardial infarction. Blood biochemical changes found after infarction have not yet been proved to have a practical value in predicting imminent ventricular fibrillation in patients with acute myocardial infarction.^{10,39}

Although an open chest procedure with general anesthesia and artificial respiration must to some extent interfere with the natural course of acute myocardial infarction these conditions allowed free choice of the site of the infarct, accurate assessment of the infarct size by use of epicardial ECG mapping and of the time of the onset of the infarct. Ideally, non-anesthetized closed chest baboons should be studied but the awake baboon can be vicious and is not readily studied in a laboratory.

If these 3 warning arrhythmias of imminent ventricular fibrillation in the baboon are applicable to patients with acute myocardial infarction the practical implications would be most important. The warning signs could detect developing ventricular electrical instability and time the onset of primary ventricular fibrillation of patients in coronary care units especially those admitted shortly after the onset of their symptoms.

To our knowledge this is the first report showing the value of warning arrhythmias

Cardiovascular events in anxiety

Experimental studies in the conscious dog

Mario Bergamaschi Ph D
Anna M Longoni D Sc
Milano Italy

During recent years attention has been increasingly focused on the possibility of controlling the emotional aspect of angina pectoris. Several minor tranquilizers—i.e. benzodiazepine derivatives have been synthesized and some of these have proved to be of help in the management of the anxious anginal patient especially when combined with long acting nitrates^{1,2}. Beta receptor blocking agents have also been used^{3,4} the usefulness of these compounds in emotional state has been based on the assumption that the autonomically mediated somatic symptoms—especially those referable to cardiovascular events—may themselves reinforce anxiety when there exists a conscious association between these symptoms and anginal pain. If so blockade of beta adrenoceptors would interrupt the somatopsychic sequence of events which are likely to be involved in the maintenance of anxiety in anginal patients.

Despite the interest in clinical cardiology little research has been reported on the cardiovascular adjustments occurring in anxiety and consequently the influence of tranquilizers and beta blockers on the hemodynamic changes brought on by this emotional state has not been defined. The majority of the reports have dealt with changes in heart rate^{5,6}. Arterial pressure^{7,8} and flow measurements⁹ have seldom been used in emotional states in man

because of technical difficulties associated with the measurements. Stebbins and Smith^{10,11} described conditioned increases in heart rate and peripheral blood flow in monkeys with ultrasonic flow probes implanted on the terminal aorta. No further studies appear to have been reported on directly measured coronary and systemic hemodynamics in emotional states.

The present paper reports the preliminary results of a study designed to investigate the influence of an anxious state induced by a classical conditioning procedure on coronary and systemic hemodynamics in trained conscious dogs; the effect of beta blockade and a minor tranquilizer (temazepam) on the altered hemodynamics was also studied.

Methods

Mongrel dogs with an average weight of 15 kilograms were used in these experiments. In general anesthesia was introduced by intravenous injection of thiopentone 50 mg per kilogram of body weight and under artificial respiration with oxygen to which halothane (2 per cent to 3 per cent) was added when required. The chest was opened through the fourth left intercostal space and the heart was exposed. The circumflex branch of the left coronary artery was dissected near its origin and an electromagnetic flow transducer was im-

From the Cardiovascular Research Institute, Milan, Italy.
Received for publication June 28, 1977.

Reprint requests to Dr. Mario Bergamaschi, Istit. C. I. Erb. Via Imbo 11/24, 20159 Milano, Italy.

- 18 Civetta J M Brons R and Gabel J C A simple and effective method of employing spontaneous positive pressure ventilation J Thorac Cardiovasc Surg 63:312 1972
- 19 Trinkle J K and Bryant L R A simple modification of existing respirators to provide constant positive pressure breathing J Thorac Cardiovasc Surg 61:617 1971
- 20 Bruyneel K and Opie L H Epicardial ECG study of the ST segment changes in the baboon with acute myocardial infarction (In preparation)
- 21 Lubbe W Bruyneel K and Opie L H Unpublished data
- 22 Brink A J Lewis C M Bosman A R and Lochner A The baboon (*Papio Ursinus*) heart (Coronary blood supply muscle function and metabolism)olia Primatol 13:11 1970
- 23 Smirk I H R waves interrupting T waves Br Heart J 11:23 1949
- 24 Carroll S L Ahuja S P and Manning G W The initiation of ventricular tachycardia and fibrillation in experimental coronary artery occlusion Am J Cardiol 16 813 1965
- 25 Lindsay A E and Budkin A The cardiac arrhythmias Chicago 1969 Year Book Medical Publishers Inc
- 26 De Sanctis R W Block P and Hutter A M Tachyarrhythmias in myocardial infarction Circulation 45 681 1972
- 27 Fisher R A and Yates I Statistical tables for biological agricultural and medical research London 1963 Oliver & Boyd Ltd
- 28 Thomas M Jewitt D E and Shillingford J P Analysis of 150 patients with acute myocardial infarction admitted to an intensive care and study unit Br Med J 1:1787 1968
- 29 Thompson P L and Stomrin G Acute myocardial infarction Predictors of arrhythmias and shock Ann Clin Res 3:377 1971
- 30 Cherry G and Myers M B The relationship to ventricular fibrillation of early tissue sodium and potassium shifts and coronary vein potassium levels in experimental myocardial infarction J Thorac Cardiovasc Surg 61:587 1971
- 31 Isomaki H Takala J and Räsänen O Influence of the site of myocardial infarction on mortality rate Acta Med Scand 185:227 1969
- 32 Thomas M Shulman G and Opie L H Arteriovenous potassium changes and ventricular arrhythmias after coronary artery occlusion Cardiovasc Res 4 327 1970
- 33 James T N Sudden death related to myocardial infarction Circulation 45:205 1972
- 34 Surawicz H Ventricular fibrillation Am J Cardiol 20 268 1971
- 35 Vatner S F Franklin D Higgins C B Patrick T White S and Van Citters R L Coronary dynamics in unrestrained conscious baboons Am J Physiol 221:1396 1971
- 36 Hill J D Mafinow M R McNulty W P and Ochsner A J Experimental myocardial infarction in unanesthetized monkeys AM HEART J 84 82 1972
- 37 Seljeskog E L Hitchcock C R Groover M E Haglin J J and Strobel C J A Isosorbide dinitrate during acute coronary occlusion Effects in the baboon JAMA 183:210 1963
- 38 Karsh R B Orlando M Norman D et al Ineffectiveness of prophylactic atropine in decreasing arrhythmias and enhancing survival following acute coronary artery occlusion in conscious dogs Am J Cardiol 29 (Abstract) 273 1972
- 39 Moss A J Goldstein S Greene W and De Camilla J Prehospital precursors of ventricular arrhythmias in acute myocardial infarction Arch Intern Med 129 756 1972
- 40 Meltzer L E and Kitchell J B The incidence of arrhythmias associated with acute myocardial infarction Progr Cardiovasc Dis 9:50 1966
- 41 Han J and Goel H G Electrophysiologic precursors of ventricular tachyarrhythmias Arch Intern Med 129:749 1972
- 42 Bennett M A and Pentecost B I Warning of cardiac arrest due to ventricular fibrillation and tachycardia Lancet 1:1351 1972
- 43 Lown B Kosowsky B and Klein M Pathogenesis prevention and treatment of arrhythmias in myocardial infarction Circulation 29:261 1969
- 44 Smirk F H and Ng J Cardiac ballet repetitions of complex electrocardiographic patterns Br Heart J 31:426 1969
- 45 Lewis T Experimental production of paroxysmal tachycardia and the effects of ligation of the coronary arteries Heart 1:98 1909
- 46 Moe G K Harris A S and Wiggers C J Analysis of the initiation of ventricular fibrillation by electrocardiographic studies Am J Physiol 134:173 1971
- 47 Harris A S and Guevara Rojas A The initiation of ventricular fibrillation due to coronary occlusion Exp Med Surg 1:105 1943
- 48 Harris A S Terminal electrocardiographic patterns in experimental anoxic coronary occlusion and hemorrhagic shock AM HEART J 35:895 1948

to 2 100 ml per minute. Blood flow through the left circumflex coronary artery ranged from 37 to 56 ml per minute with an average stroke flow of 0.58 ml. The coronary flow per unit of cardiac work was 20 ml/kg \dot{V} /min. Mean and late diastolic coronary resistance averaged 1.9 and 1.8 mm Hg/ml/min respectively. All values were fairly stable throughout the experiments.

Responses to S_1 and S_2 . The results obtained in the present experiments and the statistical evaluation of the data are reported in Table 1. Presentation of S_1 during the first experiments did not alter the cardiovascular dynamics of the trained conscious dogs to a relevant extent.

Delivery of S_2 at the fiftieth second was followed by cardiovascular changes closely resembling those described previously by Bergamaschi and co-workers¹⁰ and by Rayford and colleagues¹⁰: heart rate and cardiac output increased 108 and 120 per cent on control values; as aortic pressure increased simultaneously the averaged external work of the left ventricle rose from 2.34 kg \dot{V} per minute to 4.93 kg \dot{V} per minute. Mean coronary flow increased from 47 to 119 ml per minute and mean and late diastolic resistance decreased 33 and 40 per cent respectively. The changes in all cardiovascular parameters in the three dogs except aortic pressure were significant.

The coronary and systemic hemodynamic changes brought on by subsequent repetitions of the tone shock combination are shown in Fig. 1. Immediately after the continuous tone (S_1) was started heart rate, cardiac output, left ventricular work, and mean coronary flow increased significantly; systemic and coronary resistances showed a significant decrease. Mean aortic pressure also increased to 137 mm Hg simultaneously with the maximum increase in arterial pressure; heart rate slowed down from 202 to 126 beats per minute. Peak changes in cardiovascular dynamics were accomplished within 15 to 30 seconds. Thereafter all the cardiovascular parameters of the individual dogs decreased remaining however at levels significantly different from normals (Fig. 1 and Table 1). Heart rate was 155 beats per minute and cardiac output averaged 3.598 l per minute; vascular resistance was 38 per cent lower than control. The averaged external

work of the left ventricle was 4.9 kg \dot{V} per minute and the left circumflex coronary flow was 95 ml per minute; as both parameters were doubled during presentation of S_1 , the mean coronary flow per unit work was unchanged. Mean and late diastolic coronary resistance decreased 43 and 53 per cent respectively (Table 1).

The values of each parameter remained significantly different from controls as long as the tone was left on.

When S_2 was not presented interruption of the tone was followed by a rapid return to control values in all cardiovascular parameters (Fig. 1).

Effect of beta blockade. Propranolol 1 mg per kilogram of body weight was given intravenously at the second minute of presentation of S_1 when the pattern of cardiovascular changes induced by S_1 was fairly steady. Three minutes after administration of the beta blocker the increase in heart rate, cardiac output, left ventricular work, and mean coronary flow were markedly depressed while stroke coronary flow increased (Fig. 2).

The activity of propranolol was also investigated in a second group of trained conscious dogs in which heart rate was challenged with the infusion of increasing doses of isoprenaline. In these experiments propranolol antagonized 78 per cent the cardiac acceleration provoked by the exogenous stimulation of β adrenoreceptors (see Table 11 and Fig. 4).

Effect of the tranquilizer. Temazepam (methyl oxazepam) 10 mg per kilogram of body weight was given orally one hour before presentation of S_1 . The dose and the time were selected according to the results previously obtained in rats by Longoni, Mandelli, and Pessotti¹¹ and by Longoni and colleagues¹². The effect of the tranquilizer on systemic and coronary hemodynamic changes provoked by S_1 are shown in Table 1 and a typical tracing is given in Fig. 3. The increase in heart rate, cardiac output, and left ventricular work were depressed to levels not significantly different from controls.

Although peripheral and coronary vascular resistances increased after temazepam 10 mg per kilogram of body weight they remained significantly lower than in the quiet state. Stroke coronary flow was higher

planted on the artery, a pneumatic cuff was applied distal to the coronary flow-probe and inflation of this cuff to occlude the artery was used to determine zero flow postoperatively. A larger flowprobe was applied around the ascending aorta and an indwelling catheter was implanted in the aortic arch; the connecting wires and the catheter were brought out to the back. Experiments were not started until the dogs had completely recovered from surgery.

Aortic and coronary flow tracings were obtained by means of an electromagnetic flowmeter (Biotronix Laboratories) phasic and mean aortic pressure were measured by a Statham P23Db transducer connected to the aortic catheter; the electrocardiogram was obtained by parasternal electrodes. All signals were recorded on an 8 channel Beckman Type R Dynograph. Stroke aortic volume and stroke coronary flow were measured by planimetry of the areas beneath the phasic flow patterns.

The following parameters were calculated: mean vascular resistance, mean aortic pressure/mean flow; left diastolic (L D) coronary resistance, L D aortic pressure/L D coronary flow; vascular resistances were expressed as mm Hg/ml/min. Left ventricular work/time was calculated as a product of cardiac output (liters per minute) and mean aortic pressure (mm Hg \times 13.6) and expressed in Kg m/min.

Conditioning procedure. The experiments were carried out in daylight in a laboratory with the dogs lying on either side on a padded platform. Following the experimental model described by Estes and Skinner¹⁵ anxiety was induced in the dogs by repetitive presentations of two subsequent stimuli: S_1 , the conditioning stimulus and S_2 , the unconditioned stimulus. The former is an initially neutral stimulus which prior to training does not affect the animal's behavior to any relevant extent; the latter is a noxious stimulus which establishes the aversive drive.¹⁶ In our experiments S_1 was a continuous, low frequency tone and S_2 was a shot from a starting pistol.

During training the tone was left on for 50 seconds; at that time the shot was delivered and the tone was stopped. $S_1 \rightarrow S_2$

combination was presented to each dog once a day. On subsequent repetition the temporal relationship $S_1 \rightarrow S_2$ was progressively lengthened until S_1 alone was able to maintain the conditioned emotional responses in the recorded cardiovascular parameters.

The mean values for each parameter obtained in control state in anxiety and after treatment with temazepam were compared by analysis of variance. This was carried out following a two way cross classification treatments and dogs.

Antagonistic activity to stimulation of β adrenoreceptors. The effect of propranolol and temazepam on the cardiac acceleration provoked by isoprenaline was studied in a second group of four conscious trained dogs. Two series of increasing doses of isoprenaline were infused intravenously with a two hour interval between them. Each animal received propranolol and temazepam on separate days. Propranolol 100 μ g per kilogram of body weight was injected intravenously 15 minutes before the second isoprenaline infusion and temazepam was administered orally 1 hour before the second series of infusions.

Log dose response curves to isoprenaline before and after the two drugs were calculated by the least squares method and the distance between the two curves (Δ) was estimated on the abscissa. Δ values for propranolol and temazepam were submitted to a two way analysis of variance treatments and dogs. The potency ratios (P R) of isoprenaline were calculated as antilog of Δ values for propranolol and temazepam.

Results

Control data. Before the conditioning was started each dog was trained to lie quietly on either side on the padded platform while observations were being made. To avoid stress to the animals the same people handled the dogs throughout. Once trained all the dogs gave hemodynamic values similar to those previously reported by Gregg and co workers¹⁷ and by Bergamaschi and colleagues.¹⁸ Individual dogs had different heart rates which were in the range of 68 to 93 beats per minute. The average value of mean aortic pressure was 88 mm Hg and cardiac output ranged from 1.800

1 me 86
1 mb r 3

Table 1 Averaged results from 3 conscious dogs obtained in resting quiet state (Normal) during presentation of the conditioning stimulus S_1 before (Anxious), and 1 hour after oral administration of temazepam 10 mg per kilogram of body weight (Anxious + temazepam)

| Experiment | No dogs | Heart rate beats/min | Aortic blood pressure mm Hg | Cardiac output ml/min | Total peripheral resistance mm Hg/ml/min | Left ventricle work kg-m/min | Coronary flow | | Coronary resistance | |
|-------------------------|---------|-------------------------|--------------------------------|--------------------------|---|---------------------------------|----------------|----------------|------------------------|----------------|
| | | | | | | | Mean ml/min | Stroke ml | Vessel mm Hg/ml/min | Load dist. |
| Normal (N) | 3 | 82 ± 7.23 | 58 ± 3.05 | 1980 ± 63.40 | 0.045 ± 0.003 | 2.34 ± 0.04 | 47.5 ± 3.51 | 0.53 ± 0.07 | 1.90 ± 0.19 | 1.50 ± 0.29 |
| Anxious (A) | 3 | 155 ± 13.56 | 101 ± 1.08 | 596 ± 133.14 | 0.09 ± 0.007 | 4.93 ± 0.57 | 95 ± 15.59 | 0.60 ± 0.09 | 1.08 ± 0.07 | 0.85 ± 0.09 |
| Anxious + temazepam (T) | 3 | 90 ± 2.49 | 59 ± 2.94 | 2314 ± 114.50 | 0.039 ± 0.003 | 2.79 ± 0.07 | 0 ± 3.1 | 0.77 ± 0.05 | 1.77 ± 0.03 | 1.07 ± 0.07 |
| N → A | P | <0.01 | >0.05 | <0.01 | <0.01 | <0.01 | <0.01 | >0.05 | <0.01 | <0.01 |
| A → T | P | <0.01 | >0.05 | <0.01 | <0.01 | <0.05 | >0.05 | >0.05 | >0.05 | >0.05 |
| T → N | P | >0.05 | >0.05 | >0.05 | <0.05 | >0.05 | >0.05 | <0.05 | <0.01 | <0.03 |

The values in the table are the means with standard errors of the means and P values.

than that recorded at the same time in control experiments and the coronary flow per unit work was significantly increased from 19 ± 1.70 to 25 ± 1.77 ml per kilogram per minute ($P < 0.05$).

Temazepam 10 mg per kilogram of body weight administered by oral route 1 hour before isoprenaline did not alter the heart rate changes induced by the intravenous infusion of isoprenaline (see Table II and Fig. 4).

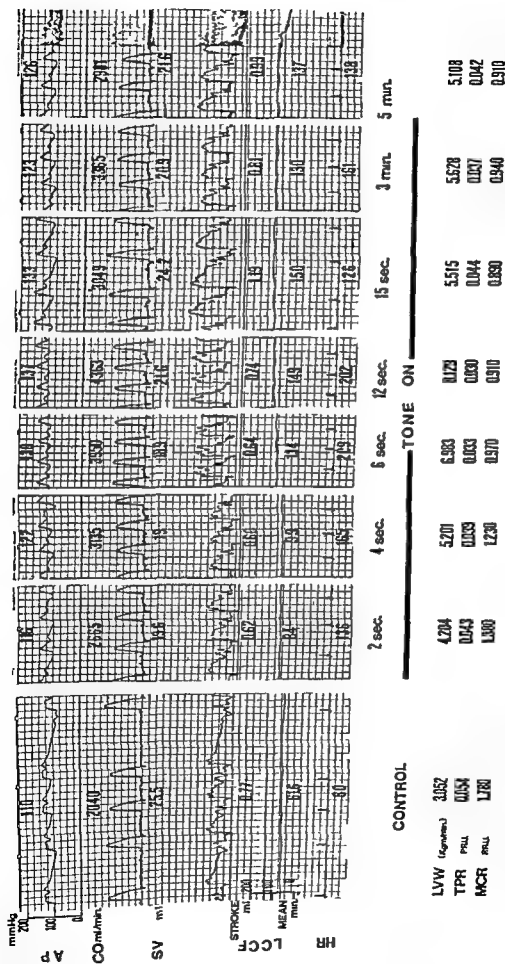
Discussion

In the classical conditioning procedure anxiety is the anticipatory response to an unpleasant event set up by presentation of an initially neutral stimulus (S_1) which was followed in the past by the aversive stimulus (S_2).¹³

The changes in heart rate, arterial pressure and peripheral flow that occur in this emotional state which somewhat resembles fear are associated with activation of the autonomic nervous system and centrally evoked somatic motor activity—i.e. uncontrolled respiration, tremor and movements of the head. These observations lead us to suggest that the anticipatory changes in the cardiovascular system are likely to be determined both by the increased sympathetic drive to the heart and vasculature and the increased oxygen demand from the muscles.¹ In order to avoid the possible

influence of the increased muscular metabolism on cardiac and vascular dynamics the dogs used in the present experiments were trained to lie still, avoiding gross muscular movements during aversive stimulation. We therefore omitted any operative response. Consequent to training, basal levels of the recorded cardiovascular parameters in the dogs were satisfactorily low. This agrees with the results obtained in previous studies in man² and in resting conscious dogs^{10, 14} which showed that sympathetic activity to the heart and vessels is minimal in the resting state in the supine position.

Delivery of the aversive stimulus (S_2) was followed by marked cardiovascular changes: heart rate, cardiac output, left ventricular work and blood flow through the left circumflex coronary artery increased significantly; peripheral and coronary resistances were markedly depressed; the changes in aortic pressure were not significant. The cardiovascular response to the aversive stimulus (S_2) was relatively constant on separate days and was always accompanied by increased respiratory activity and by movement of the head. The hemodynamic changes brought on by S_2 in the present experiments are similar to those evoked by stress in the conscious dog¹⁰ which are mediated through activation of the sympathetic nervous system as they were largely depressed by blockade of



1 Sections from a continuous record in a trained conscious dog lying on its right side showing the changes in systemic and coronary hemodynamics brought by presentation of a conditioning stimulus (tone). At the 5th minute the tone is stopped. AP = aortic pressure. CO = cardiac output. SV = stroke volume. CF = phasic (stroke) and mean (mean) blood flow through the left circumflex coronary artery. HR = heart rate and electrocardiogram.

Table II Analysis of variance of the M values (see text) for propranolol and temazepam and potency ratios (P/R) of isoprenaline before and after administration of the two drugs

| Source of variation | df | Mean square | F | I |
|---|----|-------------|-------|-------|
| Treatments (propranolol \rightarrow temazepam) | 1 | 0.9503 | 19.15 | <0.05 |
| Dogs | 3 | 0.0256 | <1 | >0.05 |
| Residual | 3 | 0.0496 | — | — |

$$P/R \frac{\text{isoprenaline} + \text{propranolol}}{\text{isoprenaline alone}} = 0.217$$

(fiducial limits for $P = 0.05$ 0.086–0.549)

$$P/R \frac{\text{isoprenaline} + \text{temazepam}}{\text{isoprenaline alone}} = 1.063$$

(fiducial limits for $I = 0.05$ 0.7202–1.569)

edly depressed the anticipatory changes in the cardiovascular system the increases in heart rate, cardiac output, left ventricular work and coronary flow were reduced. The effect on coronary flow was rate dependent as stroke coronary flow increased after propranolol. The results obtained with propranolol in the present experiments are further evidence that the conditioned changes in the cardiovascular system are largely mediated by activation of the sympathetic nervous system and adrenergic beta adrenoceptors in the heart and vessels initiated by activation of the regulatory structures in the limbic system.^{13,21} We cannot answer the question whether the efficacy of propranolol in blocking the somatic symptoms of anxiety is due to a direct action on the central nervous system or to a peripheral action on the beta adrenoceptors in the cardiovascular system. An attempt to solve the question was made by Bonn Turner and Hicks⁶ who reported that autonomically mediated somatic symptoms of anxiety in animal patients were improved after prazosin 200 mg twice daily. As very little prazosin enters the central nervous system it seems unlikely that its efficacy is related to a direct central action. A peripheral beta adrenoceptor blocking action is more probable.

Oral administration of temazepam, a benzodiazepine derivative, prevented the onset of anticipatory reactions in the cardiovascular system evoked by anxiety. The

increases in heart rate, cardiac output and left ventricular work were largely depressed by the drug. Somatic reactions also appeared diminished.

The observation that anticipatory reactions had returned to pretreatment levels on the following day indicates that the results obtained after temazepam were not influenced by a possible extinction of the emotional conditioned response consequent to removal of the unconditioned stimulus (S_u).

The neurophysiological mechanism by which temazepam depresses the altered cardiac and systemic hemodynamics in anxiety is likely to be similar to that obtained by Guerrero Figuerola and colleagues^{10,21} in conscious cats and monkeys using other benzodiazepine derivatives. They suggested that certain benzodiazepine derivatives depress the levels of anxiety, tension and fear possibly as the result of their ability both to activate the nervous structures involved in the rewarding system and to inhibit the structures responsible for aversive behavior.

The possibility that the effect of temazepam is mediated through a peripheral action on the sympathetic adrenergic receptors can be excluded since in the present experiments the tranquilizer did not alter the heart rate response to isoprenaline.

Although the increase in left circumflex coronary flow per minute produced by anxiety was depressed by the previous oral

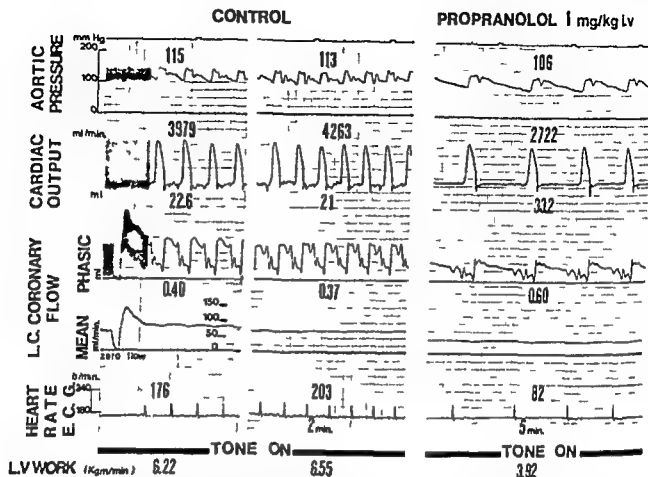


Fig 2 Effect of propranolol on anticipatory responses in the coronary and systemic hemodynamics in a dog lying on its right side. Propranolol 1 mg per kilogram of body weight was given intravenously at the second minute of the presentation of the conditioning stimulus.

the sympathetic beta adrenoceptors.¹⁹

The anticipatory reactions in the cardiovascular system evoked by the conditioning stimulus (S_1) after training were qualitatively similar to those brought on by S_2 . They started a few seconds after the onset of the tone (S_1) and reached their maximum from the fifteenth to the twentieth seconds. Reflex slowing of heart rate was commonly observed simultaneously with the peak increase in arterial pressure. During bradycardia stroke and mean coronary flow further increased while coronary resistance decreased, thus was caused by the concurrent presence of a relatively high aortic pressure, an increased duration of the diastole, and a decrease in the extravascular myocardial factors acting on the coronary vessels. All the recorded cardiovascular parameters stayed thereafter slightly below the peak effect levels remaining however, significantly different from normal values, increased respiratory activity was

always present. Similar results were obtained by Smith and Stebbins¹⁴ in conscious monkeys they reported conditioned increases in peripheral blood flow and heart rate and contended that these were true conditioned responses not secondary either to respiratory or skeletal muscle activity. The autonomic mediation of the somatic symptoms of anxiety has been demonstrated by the observation that the emotional state is associated with increased catecholamine release²⁵ and FFA mobilization.²⁶ In clinical research in anginal patients Turner and colleagues²⁴ showed that autonomic mediated somatic symptoms of anxiety were largely abolished by blockade of beta adrenoceptors with D.L. propranolol while D. propranolol, which lacks beta blocking properties was ineffective.²⁷

In the present experiments blockade of adrenergic beta receptors with propranolol 1 mg per kilogram of body weight mark

Cerebrovascular accident with unusual electrocardiographic changes

Gary J. Anderson MD
Robert Woodburn MD*
Charles Fusch MD FACC
Indianapolis Ind

Recent reports^{1,2} and reviews^{3,4} have drawn attention to the electrocardiographic (ECG) changes in cerebrovascular disease. The purpose of this case report is to describe interesting and relatively unusual electrocardiographic changes in a ten year old girl with pathologically proven intracerebral hemorrhage.

Case report

The patient (a ten year old girl) was admitted on June 19 1971 because of a left hemiparesis. The child had previously been in excellent health until the morning of admission when upon rising she complained of abdominal pain pain over her right eye and shortly thereafter developed left hemiparesis. No seizure activity was noted nor was a history of trauma elicited.

Physical examination revealed a well-developed poorly nourished female child. The apical heart rate was 60 and regular and the blood pressure was 110/90 mm Hg. Respirations were 20 per minute. Cardiac auscultation was unremarkable except for the regular rhythm. Neurological examination revealed normal R > L and both pupils showed a sluggish light response. Dysconjugate gaze was noted. Ophthalmoscopic examination was un-

remarkable. Left hemiplegia was apparent and was associated with a positive Babinski sign on the left.

Complete blood count serum sodium chloride CO₂ content blood urea nitrogen calcium and urinalyses were all normal. Serum potassium was 2.8 mEq/L. Arterial oxygen saturation was 96.6 per cent and PCO₂ was 40.0 mm Hg. Arterial pH was 7.308. Sickie cell preparation was negative. Chest x-ray and skull films were unremarkable. Cerebral angiograms were within normal limits. Lumbar puncture revealed grossly bloody cerebrospinal fluid. A rhythm strip obtained in the emergency room (Fig 1 top) demonstrated Mobitz Type I second degree AV block with junctional escape beats and periods of AV dissociation. The patient was given 0.3 mg of atropine intravenously and 1:1 conduction resumed with sinus tachycardia and wandering atrial pacemaker (Fig 1 bottom). The child's clinical course deteriorated with deepening coma and hypothermia (rectal temperature 94°F). A 12 lead ECG at that time (Fig 2) revealed sinus bradycardia with a rate of 50 per minute and a P-R interval of 0.18 second. The ST segment was elevated. Maximal ST segment elevation occurred in Leads V₂, V₃ and V₄ with an elevation exceeding 5 mm. The QT interval was 0.64 second and QT was 0.58 second. The patient died 100 hours after admission.

Necropsy examination was unremarkable except for the findings in the central nervous system.

From The Children's Hospital of Indianapolis, Indiana, and the Department of Medicine, Indiana University School of Medicine, Indianapolis, Indiana.
*Presently at the Indiana University School of Medicine, Indianapolis, Indiana.
Reprint requests: Gary J. Anderson MD, 1100 West Michigan Street, Indianapolis, Indiana 46202.
Dr. Anderson is a resident in the Department of Medicine, Indiana University School of Medicine, Indianapolis, Indiana.

- sure changes conditioned to painful stimuli (schizokinesis) *Bull Johns Hopkins Hosp* 107:72 1960
- 11 Katcher A H Solomon R L Turner L H Lo Lordo V Overmier J B and Rescorla R A Heart rate and blood pressure responses to signaled and unsignaled shock effect of cardiac sympathectomy *J Comp Physiol Psychol* 68:163 1969
 - 12 Shmavonian H M Methodological study of vasomotor conditioning in human subjects *J Comp Physiol Psychol* 52:315 1959
 - 13 Stebbins W C and Smith O A Cardiovascular concomitants of the conditioned emotional responses in the monkeys *Science* 144:881 1964
 - 14 Smith O A and Stebbins W C Conditioned blood flow and heart rate in monkeys *J Comp Physiol Psychol* 59:432 1965
 - 15 Estes W K and Skinner B F Some quantitative properties of anxiety *J Exp Psychol* 29:390 1941
 - 16 Schoenfeld W N An experimental approach to anxiety escape and avoidance behaviour in Hoch P H and Zubin J editors *Anxiety* New York 1964 Haffner Publishing Co p 70
 - 17 Gregg D E Khouri E M and Rayford C R Systemic and coronary energetics in the resting unanesthetized dog *Circ Res* 16:102 1965
 - 18 Bergamaschi M Shanks R G Caravaggi A M and Mandelli V A comparison of the cardiovascular actions of four adrenergic β receptor blocking agents in resting conscious dogs *Am Heart J* 82:338 1971
 - 19 Bergamaschi M Caravaggi A M Mandelli V and Shanks R G Sympathetic control of coronary and systemic hemodynamics in emotional stress in conscious dogs *Am Heart J* (In press)
 - 20 Rayford C R Khouri E M and Gregg D E Effect of excitement on coronary and systemic energetics in unanesthetized dogs *Am J Physiol* 209:680 1965
 - 21 Longoni A Mandelli V and Pessotti I Variable interval in rats treated with temazepam diazepam oxazepam and chlordinzepam *Pharmacol Res Comm* 3:165 1971
 - 22 Longoni A Mandelli V and Pessotti I Conditioned suppression in rats treated with temazepam (In press)
 - 23 Robinson B F Epstein S E Berser G D and Braunwald D E Control of heart rate by the autonomic nervous system *Circ Res* 19:400 1966
 - 24 Pitt B Greene H L Sugishita Y and Ross R S Effect of β adrenergic receptor blockade on coronary hemodynamics in the supine anesthetized dog *Cardiovasc Res* 4:109 1970
 - 25 Elmudjan F Hope J M and Lamson E T Excretion of epinephrine and norepinephrine in various emotional states *J Clin Endocrinol Metab* 17:608 1957
 - 26 Vouridis E Trichopoulos A Cokkinos D Plessas S T and Miras C Free fatty acid elevation during diagnostic cardiac procedures Communication at the VI European Congress of Cardiology Madrid 23-30 September 1972
 - 27 Bonn J A and Turner P D propranolol and anxiety *Lancet* II:1355 1971
 - 28 Nauta W J H Limbic system and hypothalamus anatomical aspects *Physiol Rev* 40 (Suppl 4) 102 1960
 - 29 Hilton S M and Zbrozyna A W Amygdaloid region for defence reactions and its afferent pathway to the brain stem *J Physiol* 165:160 1963
 - 30 Guerrero-Figueroa R Rye M M and Heath R G Effects of two benzodiazepine derivatives on cortical and subcortical epileptogenic tissues in the cat and monkey I Limbic structures *Curr Ther Res* 11:27 1969
 - 31 Guerrero-Figueroa R and Gallant D M Electrophysiological study of the action of a new benzodiazepine derivative (ORF 8063) on the central nervous system *Curr Ther Res* 13:747 1971
 - 32 Abel R M Reis R L and Starosck R N Coronary vasodilatation following diazepam (Valium) *Br J Pharmacol* 38:620 1970
 - 33 Abel R M Reis R L and Starosck R N The pharmacological basis of coronary and systemic vasodilator actions of diazepam (Valium) *Br J Pharmacol* 39:261 1970
 - 34 Bergamaschi M Fucella L M Mandelli V Tommasini R Turba C and Usardi M M Pharmacological and clinical studies on a new beta adrenergic blocking agent the 1-(1-isopropylamino-3-(1,2,3,4-tetrahydro-1,4-ethano 5-naphthyl)oxy)-2-propanol HCl (K 4123) Nonyl Schmiedebergs *Arch Pharmacol* 269:447 1971

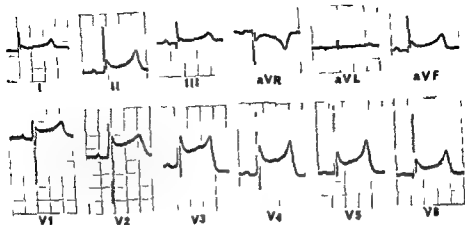


Fig. 2 Twelve-lead electrocardiogram 74 hours after admission (see text)

that intracerebral disease may induce subendocardial hemorrhage⁸ and ultrastructural myocardial changes.^{2,22} Such changes may be of sufficient magnitude to induce Q waves.⁸ While the association between cerebrovascular accidents and myocardial infarction has been made in patients prone to both conditions²³ the question may be raised as to whether ST segment elevation may be due to myoelectrolysis alone. The crick reported demonstrated normal microscopic examination of the heart suggesting that the ST segment changes were mediated either by alteration of autonomic tone pH electrolyte disturbances or a combination of all of these factors.

Summary

This case report describes a 10-year-old girl sustaining a cerebrovascular accident secondary to a vascular malformation involving the cingulate gyrus. The patient developed a Mobitz Type I block with periods of AV dissociation. A 12-lead electrocardiogram revealed the frequently observed prolongation of the QT interval. Her ECG also demonstrated diffuse ST segment elevation. Neurophysiological mechanisms are discussed.

REFERENCES

1. Birch G E, Myers R and Wilder J A. A new electrocardiographic pattern observed in cerebrovascular accidents. *Circulation* 19:719 1954.
2. Levin H. Non-specificity of the electrocardiogram associated with coronary artery disease. *Am J Med* 15:344 1953.
3. Hersch C. Electrocardiographic changes in head injury. *Circulation* 28:853 1961.
4. Hersch C. Electrocardiographic changes in subarachnoid hemorrhage meningitis and intracranial space-occupying lesions. *Br Heart J* 26:785 1964.
5. Wilder J A, Miller K, Burgess M J and Vincent W. The electrocardiogram and the central nervous system. *Progr Cardiovasc Dis* 13:210 1970.
6. Greenhoot J H and Reichenbach H D. Cardiac injury and subarachnoid hemorrhage: a clinical, pathological and physiological correlation. *J Neurosurg* 30:521 1969.
7. Wilder J A. Electrocardiographic wave forms and the nervous system. *Circulation* 41:371 1970.
8. Kortebein G C, Boles J F and Ten Cate J. Influence of stimulation of some subcortical areas on electrocardiogram. *J Neurophysiol* 20:100 1957.
9. Fulton J F. Functional localization in the frontal lobes and cerebellum. London 1949. Oxford University Press p 66.
10. Jacobson S A and Danoff J. Marked electrocardiographic changes produced by head trauma. *J Neuropathol Exp Neurol* 13:462 1954.
11. Smith M and Ray C T. Cardiac arrhythmia: increased intracranial pressure and the autonomic nervous system. *Dis Chest* 61:125 1972.
12. Mauch H I, Hockman C H and Hoff E C. ECG changes after cerebral stimulation I. Anomalous atrioventricular excitation elicited by electrical stimulation of the mesencephalic reticular formation. *Am Heart J* 38:198 1964.
13. Loole J L. Vasocardiac effects of the circle of Willis. *Arch Neurol Psychiatr* 8:355 1957.
14. Brodal A. Neurological anatomy in relation to clinical medicine. London 1969. Oxford University Press p 662.
15. Bearl F I, Robertson J W and Robertson R C. Spontaneous subarachnoid hemorrhage stimulating acute myocardial infarction. *Am Heart J* 8:735 1959.

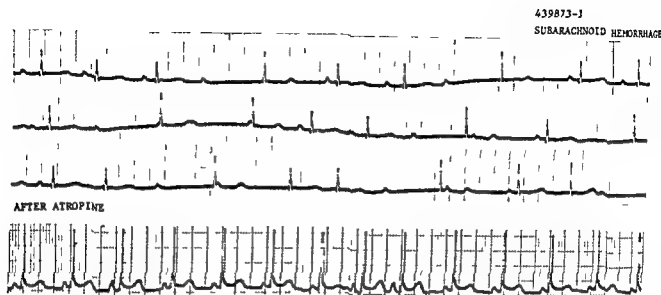


Fig. 1 Top panel Rhythm strip of the patient on admission demonstrating Mobitz Type I AV block with periods of AV dissociation. Bottom panel Restoration of 1:1 conduction with wandering, atrial premature after 0.3 mg atropine intravenously.

A vascular malformation was found to involve the cingulate gyrus at the right frontal parietal junction. This vascular malformation had ruptured resulting in intraventricular and subarachnoid hemorrhage. Transient and transfrontal herniation had occurred.

Discussion

The case reported is characterized by two interesting electrocardiographic changes.

Fig. 1 demonstrates second degree AV block and periods of AV dissociation with return of AV conduction to normal following administration of atropine. To our knowledge the Wenckebach block seen in this patient clearly related to a cerebrovascular accident has not been previously described although 2:1 block has been observed.⁴

Recent papers^{8,11} have drawn attention to the fact that increased vagal tone contributes to the development of the observed arrhythmias in cerebrovascular accident. Experimental studies¹² have shown that the production of Wolff Parkinson White pre-excitation and AV dissociation¹² may occur with electrical stimulation of the mesencephalic reticular formation or the dorsomedial hypothalamic nucleus. These experimental studies suggest that enhanced vagal tone significantly contributes to the development of various cardiac arrhythmias. Such enhanced vagal tone may be a result of stimulation of vagal fibers innervating the circle of Willis¹³ enhanced vagal

tone secondary to the baroreceptor reflex⁴ or stimulation of Area 13, the chief cortical representation of the vagus nerve.⁵ In our patient the lesion involved the cingulate gyrus containing Areas 23, 24, and 37. These areas are thought to control autonomic functions of the heart.¹⁴ Thus involvement of the cingulate gyrus may have accounted for the observed cardiac arrhythmia.

A second interesting feature of this case is the S1 and T wave changes. Cerebrovascular accidents are as a rule associated with S1 segment prolongation and depression¹ and T wave inversion,^{1,2} occasionally simulating myocardial infarction.^{1,15} On the other hand S1 segment elevation is less frequently observed.¹⁶ It has been proposed¹ that such changes may be secondary to electrolyte imbalance, pH and blood gas abnormalities, the latter supported by experimental studies.^{17,18} Abnormalities of serum K⁺ and pH were observed in the reported case and may have contributed to the LCF pattern of elevated S1 segment.¹⁹ Hypoalbumin has also been implicated in the development of second degree heart block.¹⁹ The second positive deflection at the J junction (Fig. 2 leads II, III, aVR and V₁ through V₆) as well as the prolonged QT may have been due in part to hypothermia^{20,21} in addition to enhanced vagal tone.

Experimental studies have related

- 16 Cropp G J and Manning G J Electrocardiographic changes simulating myocardial ischemia and infarction associated with spontaneous intracranial hemorrhage *Circulation* 22:25 1960
- 17 Roberts K E and Magida M G Electrocardiographic alterations produced by a decrease in plasma pH bicarbonate and sodium as compared with those produced by an increase in potassium *Circ Res* 1:206 1953
- 18 Magida M G and Roberts K E Electrocardiographic alterations produced by an increase in plasma pH bicarbonate and sodium as compared with those seen in a decrease in potassium *Circ Res* 1:214 1953
- 19 Fisch C Knoebel S B Leikenbaum H and Greenspan K Potassium and the monophasic action potential electrocardiogram conduction and arrhythmias *Progr Cardiovasc Dis* 8:387 1966
- 20 Osborn J J Experimental hypothermia respiratory and blood pH changes in relation to cardiac function *Am J Physiol* 116:389 1953
- 21 Byer E Ashman R and Toter L A Electrocardiogram with large upright T waves and long QT intervals *AM HEART J* 33:96 1947
- 22 Burch G E Sobral R S Sun S C and Colcolough H L Effects of experimental intracranial hemorrhage on the ultrastructure of the myocardium of mice *AM HEART J* 17:427 1969
- 23 Connor R C Heart damage associated with intracranial lesions *Br Med J* 3:79 1968

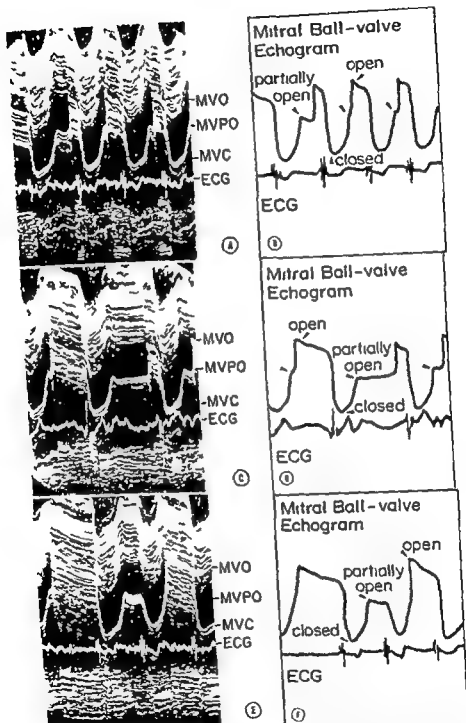


Fig 2 A through F Echocardiograms of mitral ball valve prosthesis. MVO = ball in open position. MVPO = ball in partially open position. MVC = ball in closed position. On the left Polaroid photographs of echocardiograms (A, C and E) and on the right (B, D and F) the corresponding artist's line drawings of original recording of the malfunctioning Cutter Smeloff mitral valve prosthesis. The echocardiograms illustrate the varying degree of valve dysfunction, particularly the tendency for the ball to stick in the partially open position during part of diastole with the result that the time spent in the partially open position frequently varies from beat to beat. In the middle beat of E and F the ball never reaches the fully open position. It should be noted that since the echo beam seldom if ever directed precisely along the pathway of the moving ball different views of the ball are probably recorded during the cardiac cycle.

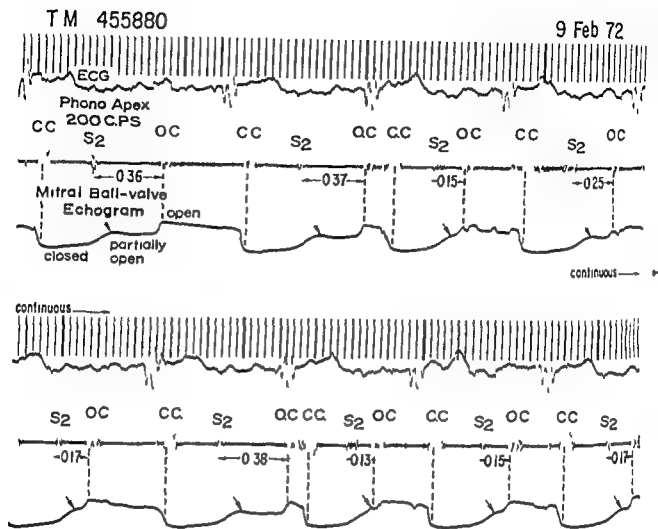


Fig 1 Simultaneous echocardiography and phonocardiography. Phono = phonocardiogram taken at the apex at 200 cycles per second. CC = closing click. OC = opening click. S_2 = second heart sound. S_2 -OC intervals are shown in seconds (varying from 0.13 to 0.38). The points at which the ball begins to stick in the partially open position are shown by arrows. The OCs correspond with the arrival of the ball to the fully open position. The CC corresponds to the ball moving to the closed position. The echocardiograms which were recorded in the analog method show a rather consistent location of the maximal anterior movement of the ball coincidental with the opening click. This consistency supports the conclusion that the points of the maximal excursion are real rather than artifactual.

On February 15, 1977, after three weeks of intensive antimicrobial therapy for possible bacterial endocarditis, the patient was operated on and the prosthesis was replaced with a Ballie die valve mitral prosthesis. The left atrium was markedly enlarged and its posterior wall between the insertion of the left and right pulmonary veins was completely covered with old and recent thrombi which were removed. The orifice of the mitral valve was narrowed by the overgrown fibrous tissue over the valvular ring. The struts of the valve (Fig. 3) were segmentally covered with various size of similar fibrous tissue which impeded the movement of the ball both in the closed position and as the ball passed through the ring of the prosthesis. No bacteria were seen microscopically or in cultures obtained at surgery.

Comments

Malfunction of a mitral ball valve prosthesis is not commonly recognized but

when present is associated with symptoms due to cerebral emboli and/or congestive heart failure.⁴ In several of the cases reported, repeated neurological symptoms occurred often months or several years (as in this patient) before the prostheses were eventually replaced. Since peripheral embolization and congestive failure frequently occur in patients with rheumatic valvular disease with large left atria and atrial fibrillation, their presence does not necessarily implicate a malfunctioning valve as their cause. Because of the serious morbidity often associated with systemic emboli and heart failure, however, the early detection of a malfunctioning prosthetic valve is of obvious importance.

Malfunction of a mitral

echocardiographically. It would appear reasonable to perform these studies orally at intervals of 3 to 6 months in patients with a mitral ball valve prosthesis and whenever symptoms suggestive of peripheral embolization or of the onset or worsening of congestive heart failure are present.

REFERENCES

- 1 Hylen J C, Kloster F E, Herr R H, Starr A and Griswold H F. Phonocardiographic diagnosis of aortic ball variance. *Circulation* 38:90 1968.
- 2 Hylen J C, Kloster F E, Herr R H, Starr A and Griswold H E. Sound spectrographic diagnosis of aortic ball variance. *Circulation* 39:849 1969.
- 3 Hylen J C, Kloster F E, Starr A and Griswold H E. Aortic ball variance diagnosis and treatment. *Ann Intern Med* 74:1 1970.
- 4 Sander on R G, Hall A D and Thomas A N. The clinical diagnosis of ball variance in a mitral valve prosthesis. *Ann Thorac Surg* 6:473 1968.
- 5 Leatherman L L, Leachman R D, McConn R G, Halimin G L and Cooley D A. Malfunction of mitral ball valve prosthesis due to swollen poppet. *J Thorac Cardiovasc Surg* 57:160 1969.
- 6 Lee S J, K Zargoza A J, Callaghan J C, Couves C M and Sterns L J. Malfunction of the mitral valve prosthesis (Cutter Smeloff). *Circulation* 41:419 1970.
- 7 McHenry J M, Smeloff E A, Fong W Y, Miller C E and Ryan P M. Critical obstruction of prosthetic heart valves due to lipid absorption by silastic. *J Thorac Cardiovasc Surg* 59:413 1970.
- 8 Craig E, Hutchins P and Sutton R. Impaired function of cloth-covered Starr Edwards mitral valve prosthesis. *Circulation* 41:141 1970.
- 9 Garimella J J, Lynch M I, Schmidt W R and Jensen N H. Fatal clotting of the Starr Edwards mitral ball valve nineteen months postoperatively. *J Thorac Cardiovasc Surg* 46:673 1964.
- 10 Schlager J, Minna F P and Wolf R E. Auscultatory and phonocardiographic signs of ball variance in a mitral prosthetic valve. *Am Heart J* 81:809 1971.
- 11 Pfeifer J, Goldschlager N, Sweetman T, Gerbode E and Selzer A. Malfunction of mitral ball valve prosthesis due to thrombus. *Am J Cardiol* 29:95 1972.
- 12 McHenry J M, Smeloff E A, Davey T B, Kaufman B and Fong W Y. Hemodynamic results with full flow orifice prosthetic valves. *Circulation* 34 (Suppl 1):24 1967.
- 13 Winters W L, Gimenez J L and Goloff L A. Clinical application of ultrasound in the analysis of prosthetic ball valve function. *Am J Cardiol* 19:97 1967.
- 14 Johnson M C, Lyon H C and Holmes J H. Ultrasonic evaluation of prosthetic valve motion. *Circulation* 41 and 42 (Suppl II):3 1970.
- 15 Siglers D C, Srivongse S A and Deuchar D. Analysis of dynamics of mitral Starr Edwards valve prosthesis using reflected ultrasound. *Br Heart J* 33:401 1971.
- 16 Craig M. Phonocardiographic studies in mitral stenosis. *N Engl J Med* 257:650 1957.
- 17 Zimik R S and Burchell H B. A phonocardiographic study of patients with total prosthetic mitral valve replacement. *Dis Chest* 44:11 1963.
- 18 Nijimi M and Segal B L. Auscultatory and phonocardiographic findings in patients with prosthetic ball valves. *Am J Cardiol* 16:793 1965.

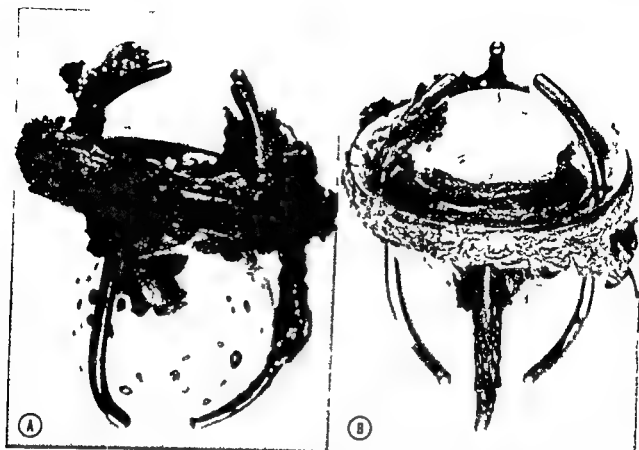


Fig 3 A and B Photographs of the Cutter Smeloff ball valve prosthesis removed at surgery in the open position (A) and in the stuck position as the ball passes through the ring (B). There is extensive fibrous overgrowth on the valve ring as well as on the atrial and ventricular struts.

thesis can be suspected by prolonged and variable intervals between the second heart sound and the opening click of the prosthetic valve which should normally be less than 0.15 second.^{8,10,12,13} In all but one of the reports cited above,⁴⁻¹¹ the intervals were recorded phonocardiographically and shown to be variable and prolonged. Occasionally the opening click was intermittently absent. In the one exception,⁹ the valve sounds were described as being normal but phonocardiography was not performed.

In atrial fibrillation the variation of the S₂ opening click intervals with preceding cycle lengths may be explained by the longer diastolic filling phase allowing the left atrium to decompress thereby lowering left atrial pressure and the left atrial to left ventricle pressure difference. (This is similar to the changes in the S₂ OS intervals in mitral stenosis.^{16,19}) The lower pressure gradient causes the S₂ opening click interval to be prolonged and as evidenced in Figs 1 and 2, the interval is also prolonged in malfunctioning prosthesis.

Echocardiography appears to be a promising and noninvasive technique for the assessment of prosthetic valve function. The motion of the prosthetic ball resembles that of the anterior leaflet of the mitral valve in mitral stenosis.^{20,21} The dimension of the ball as well as its excursion, timing and speed of motion are all easily obtained and reproducible at least over short periods of time.²² To our knowledge Pfeifer and associates²³ reported the first use of ultrasound in the detection of prosthetic ball valve malfunction. The echograms shown in their report showed that the ball stuck in the closed position whereas in the prosthesis removed from our patient there was a tendency for the ball to be stuck both in the closed posterior position and again as it passed through the ring of the prosthesis.

Summary

Mitral ball valve prosthesis malfunction is uncommon but may be associated with serious morbidity and may be fatal. The diagnosis may be suspected clinically and confirmed by phonocardiography^{4,11,24} and

- B Apparently isolated deoxycortico-sterone excess⁶⁴
- C Congenital abnormalities of corticosteroid synthesis
 - 1 17 α hydroxylation deficiency^{67, 68}
 - 2 11 β hydroxylation deficiency⁶⁹
 - 3 21 hydroxylation deficiency⁶⁸
 - 4 Hypertension associated with some cases of Turner's syndrome⁶⁴
- D Cushing's syndrome⁶
- E Corticosterone excess⁶¹
- F Licorice^{62, 66} and carbenoxolone induced hypertension^{61, 67}

LOW RENIN HYPERTENSION WHERE NO CAUSAL AGENT IS FOUND^{12, 13, 70}

- A Mineralocorticoid excess suspected^{68, 70}
- B Mineralocorticoid excess not implicated⁷¹

Low renin hyperaldosteronism

ALDOSTERONE SECRETING ADRENOCORTICAL ADENOMA This is the classical form of the disease originally described by Conn² as a benign adenoma of the adrenal cortex secreting an excess of aldosterone and giving rise to increased plasma levels of aldosterone. The effect of the elevated circulating aldosterone is to promote sodium retention and potassium wasting hence total exchangeable and plasma sodium are elevated total exchangeable and total body potassium are reduced and hypokalemia usually occurs.⁷ In association with the sodium retention total body water extracellular fluid and plasma volumes are increased^{7, 11, 6} and there is a resultant depression of circulating levels of renin and angiotensin II.^{11, 12, 16} In this syndrome therefore the aldosterone secretion is clearly not a consequence of stimulation by the renin/angiotensin system⁷² indeed in our own series a significant inverse correlation has been demonstrated between current measurements of plasma concentrations of renin and aldosterone.^{11, 15}

The hypertension appears to be closely related to the sodium retention and we have discussed elsewhere in some detail possible mechanisms underlying the development of high blood pressure.^{15, 11, 16}

Surgical removal of the adenoma leads to a fall of aldosterone secretion rate and plasma aldosterone with consequent natriuresis and correction of the hypokalemia

Plasma concentrations of renin and angiotensin II increase as the total exchangeable sodium falls and the hypertension usually, although not invariably is corrected.^{11, 15}

The effects of spironolactone in counteracting aldosterone excess and controlling the hypertension have been clearly demonstrated in this syndrome. The dose of spironolactone has to be substantial (up to 400 mg per day) in view of the often markedly elevated aldosterone secretion rate. With adequate therapy total exchangeable sodium and plasma sodium are reduced as are total body water extracellular fluid and plasma volumes. Total exchangeable potassium and total body potassium increase and plasma potassium rises into the normal range.^{11, 16, 10} Extracellular alkalosis is also corrected and plasma bicarbonate concentration falls. Plasma levels of renin and angiotensin II rise as the total exchangeable sodium falls and sometimes abnormally high levels of renin and angiotensin are observed during treatment possibly indicating over correction and slight sodium depletion. The effect of spironolactone on aldosterone secretion rate is variable. With treatment several potential stimuli to aldosterone secretion are activated (potassium retention sodium depletion and elevation of angiotensin II). Spark and Melby¹¹ found that aldosterone secretion rate increased during therapy in some cases although Brown and colleagues^{11, 74} observed that it might remain unaltered indicating relative autonomy of the adrenocortical adenoma. Plasma aldosterone concentration may be elevated during therapy¹¹ this perhaps partly due to an increase in aldosterone secretion partly to diminution in aldosterone metabolic clearance possibly related to reduction in hepatic blood flow and partly to shrinkage of plasma volume.

ADRENOCORTICAL MICROADULAR HYPERPLASIA It is now recognized that not all cases of low renin hyperaldosteronism are due to aldosterone secreting adrenocortical adenomata. In approximately 25 per cent of cases the adrenal cortices show diffuse usually bilateral micronodular hyperplasia simple hyperplasia of the zona glomerulosa or even in a small proportion no recognizable histological change.^{6, 11, 11} It is possible in these cases that the excess of

The use of spironolactone in the diagnosis and the treatment of hypertension associated with mineralocorticoid excess

D G Beevers, M R C P

J J Brown, F R C P

J B Ferriss M R C P

R Fraser, Ph D

A F Lister F R C P

J I S Robertson I R C P

Glasgow, Scotland

Spiroinolactone is a steroidal lactone which antagonizes the actions of aldosterone particularly in distal renal tubules where it promotes sodium excretion and potassium retention. The drug is frequently employed for its diuretic actions but here we will consider its use as a specific agent in correcting the hypertension and electrolyte abnormalities of hyperaldosteronism and other syndromes of mineralocorticoid excess. For this purpose the forms of hypertension where mineralocorticoid excess has been implicated have been classified below. Clearly in some of the rarer syndromes spiroinolactone therapy is not appropriate and they are merely included for completeness.

Classification of hypertension associated with excess of mineralocorticoids¹

I ALDOSTERONE EXCESS WITH LOW PLASMA RENIN

- A Aldosterone producing adrenocortical adenoma ("primary aldosteronism")²⁻⁶

- B Adrenocortical micronodular hyperplasia (idiopathic hyperaldosteronism) pseudo primary aldosteronism)⁶⁻¹²

- C Adrenocortical carcinoma¹³⁻¹⁴

- D Dexamethasone responsive hyperaldosteronism¹⁵⁻¹⁸ (Glucocorticoid remediable hyperaldosteronism)

II ALDOSTERONE EXCESS ASSOCIATED WITH A HIGH PLASMA RENIN

- A Malignant phase hypertension¹⁹⁻²²

- B Renal artery stenosis²³⁻²⁵

- C Severe bilateral chronic renal disease²⁶⁻²⁸

- D Renin secreting renal tumor²⁹⁻³¹

- E Congenital hyperaldosteronism³²⁻³⁴

- F Following treatment with benzothiadiazines³⁵⁻³⁷

III HYPERTENSION OF PREGNANCY³⁸⁻⁴¹ AND HYPERTENSION DUE TO ORAL CONTRACEPTIVE DRUGS⁴²⁻⁴⁵

IV NON ALDOSTERONIC MINERALOCORTICOID EXCESS

- A 18 Hydroxy deoxycorticosterone excess⁴⁶⁻⁴⁸

From the M R C Blood Pressure Unit Western Infirmary Glasgow Scotland
Received for publication Dec 8 1972

Reprint requests to Mrs A McGregor M R C Blood Pressure Unit Western Infirmary Glasgow G11 6NT Scotland

is more likely in patients with evidence of renal impairment before starting treatment^{11 10}

ADRENOCORTICAL CARCINOMA Occasional cases of hyperaldosteronism with hypertension due to a carcinoma of the adrenal cortex have been described^{12 24} It is important that this is borne in mind before embarking on long term spironolactone therapy Most examples of carcinoma have had features which should readily distinguish them from adenomata The tumors have usually been large frequently displacing the kidney on intravenous urography or arteriography fever and abdominal pain have been common and usually other corticosteroids in addition to aldosterone have been produced in excess²⁵

DEXAMETHASONE RESPONSIVE ALDOSTERONISM This syndrome is probably extremely rare^{26 27} and is usually associated with hyperplasia or nodular changes in the adrenal cortex It is our usual practice before starting spironolactone to test the effect of dexamethasone in a dose of 2 mg per day for 2 weeks To date we have not seen an example of dexamethasone responsiveness

Secondary hyperaldosteronism

These syndromes are seen in association with unilateral or bilateral renal or renal artery disease^{1 20 21 28 29} and in malignant phase hypertension^{1 30 31} with or without an identifiable cause An important variety is seen in those patients with terminal renal failure treated with hemodialysis whose blood pressure is not readily controlled The hyperaldosteronism is associated with elevated levels of renin and angiotensin and in contrast to primary aldosteronism significant positive correlations of concurrent measurements of renin angiotensin I and II and aldosterone have been demonstrated^{25 26 27 32 33}

RENAL ARTERY STENOSIS In severe cases of renal artery stenosis where the malignant phase is usually present^{34 35 36} patients may have markedly elevated levels of renin angiotensin II and aldosterone and an associated reduction in plasma and total exchangeable potassium However in contrast to low renin hyperaldosteronism the plasma and total exchangeable sodium may be reduced^{31 32 33} Surgical relief of the stenosis may lead to correction of the hypertension and the biochemical abnor-

malities Clearly the hypertension is not simply due to the sodium and water retention of aldosterone excess²⁸ and it is postulated that the reduction in blood flow to the affected kidney causes an intense signal for renin release inappropriate to the requirements of the body The resultant high levels of angiotensin II stimulate aldosterone secretion which favors sodium retention and elevation of the blood pressure However the raised arterial pressure provokes natriuresis from the unaffected kidney and the increased levels of angiotensin II cause sodium loss from both the poststenotic and unaffected sides³⁷ The overall effect appears to be a slight reduction in the total exchangeable sodium It is possible that in these circumstances the elevated levels of angiotensin II are sufficient to have a direct pressor effect However in many less severe cases of renal artery stenosis with hypertension³⁸ the peripheral plasma concentrations of renin and angiotensin II are within the normal range³⁹ Even in these however it is possible that renin and angiotensin II are inappropriately high in relation to sodium status^{41 42}

MALIGNANT PHASE HYPERTENSION A very similar syndrome may be seen in malignant phase hypertension in the absence of a demonstrable renal or renal artery lesion^{1 20 21} It has been suggested that in this situation intrarenal arterial and arteriolar lesions resulting from the malignant phase may produce a similar effect on renin secretion to that of renal artery stenosis⁴³ Anti-hypertensive therapy permits the histological lesions to heal and the elevated levels of renin angiotensin II and aldosterone revert to normal

INTRACTABLE HYPERTENSION IN TERMINAL RENAL FAILURE In the majority of patients undergoing regular hemodialysis for chronic renal failure blood pressure is readily controlled by removal of sodium and water at dialysis with restriction of sodium and water intake between dialyses In a minority however these measures are ineffective and severe intractable hypertension develops in association with very high levels of circulating renin angiotensins I and II and aldosterone^{44 45 46} Removal of salt and water at dialysis fails to control the blood pressure resulting merely in further eleva-

aldosterone is under the stimulus of some as yet unidentified trophic factor

Spark and Melby³⁰ found spironolactone to be ineffective in controlling the hypertension in two cases without adenoma, and suggested that such lack of response might be general in this variant of the disease and thus be the basis of a preoperative test, differentiating the adenoma and non adenoma groups. Crane and Harris³² however, obtained a good response to spironolactone in two non adenomatous patients and Brown and colleagues³¹ in a more extensive series found no significant difference in the hypotensive response to prolonged spironolactone therapy in 21 patients with and 11 without adrenocortical adenoma. In both adenomatous and non adenomatous groups occasional lack of hypotensive response was usually associated with evidence of renal impairment before the start of therapy. This experience did, however, confirm Spark and Melby's finding³⁰ that the hypotensive response to preoperative spironolactone usually predicted very closely the subsequent effect of adrenocortical surgery.

It is our practice to give spironolactone to within 24 hours of operation thus presenting the patient to surgery with normal electrolyte status. Furthermore, preoperative spironolactone may combat or minimize the development of postoperative hypoaldosteronism.³³ It is emphasized that spironolactone or its breakdown products give a fluorescence in plasma very similar to that of cortisol; if therefore the Mattingsly test³⁴ is used to measure 11 hydroxy corticosteroids, a falsely high result will be given if the patient is receiving spironolactone.^{1, 35, 36} This effect takes some 14 days to clear after therapy is withdrawn.

Although the response to spironolactone cannot be used in distinguishing cases with from those without adrenocortical adenoma, this distinction can be made on biochemical grounds. Several workers have observed that the biochemical features of the disease are generally more marked in the adenoma group.^{3, 10, 31} plasma levels of aldosterone, sodium, and bicarbonate tending to be higher and renin and potassium lower. Individually, these measurements are not usually sufficient to provide a firm basis for diagnosis, but a multifac-

torial computer assisted technique evaluating them all simultaneously (quadruc analysis) has been successfully and consistently utilized in distinguishing patients with from those without adrenocortical adenoma.³¹

Prolonged therapy with large doses of spironolactone has been shown to be effective in a large series of patients with and without adrenocortical adenoma¹² with no tendency for the blood pressure to escape from control even after periods of up to 9 years. Thus spironolactone treatment can now be offered as a reasonable alternative to the extensive bilateral adrenocortical resection otherwise required in patients shown on the basis of quadruc analysis to be unlikely to have an adrenocortical adenoma. Spironolactone can similarly be employed in patients unwilling or unsuitable to undergo operation. Although the drug has many possible side effects in our experience these are not usually severe and in only 2 of 67 cases was it necessary to withdraw therapy.¹¹ This contrasts with earlier statements³⁰ that side effects usually rendered long term therapy intolerable. Epigastric discomfort following ingestion of the tablets can usually be minimized if they are taken with or after food although spironolactone is inadvisable in patients with peptic ulceration. Gynecomastia occurs frequently in male patients after several months of therapy in high dosage³⁴ there have been occasional complaints of impotence. In women menorrhagia, oligomenorrhea or occasional amenorrhea occurs. Other complaints have been constipation, excessive sweating, lassitude (especially in the early months of therapy), alopecia, Rignaud's phenomenon and cutaneous pigmentation. Patients unresponsive to both surgery and spironolactone generally have evidence of renal impairment a finding which is in accordance with the notion that perpetuation of the hypertension might be related to intra renal vascular damage consequent upon the raised arterial pressure.³⁷

Spironolactone irrespective of the hypotensive effect invariably corrects the abnormal electrolyte pattern; consequently potassium supplements are not required and may indeed be dangerous. Elevation of plasma potassium to abnormally high levels

J. I. me 86
A. mb 3

droxycorticosteroids by the Mattingly technique^{1, 2, 3, 4}

In one case of corticosterone excess associated with an adrenocortical carcinoma in a patient with hypertension hypokalemia hypernatremia and peripheral edema a 6 day course of spironolactone 300 mg daily was ineffective.¹¹

Licorice extracts contain the ammonium salt of glycyrrhizic acid a substance with an aldosterone like action. As a result persons who consume large quantities of licorice-containing sweets and drinks may develop hypertension with hypernatremia low plasma renin and hypokalemic alkalosis.¹²⁻¹⁶ This syndrome may be corrected by spironolactone treatment.¹⁴ A similar clinical picture may be induced by therapy with carbenoxolone^{1, 17} but spironolactone is contraindicated in its correction as it also blocks the ulcer healing effect.

Low renin hypertension

It is now apparent that approximately 25 per cent of patients with apparently essential hypertension have low levels of plasma renin activity.¹⁸⁻²² It has been postulated that the suppression of renin in many of these patients may be related to excess of an as yet unidentified mineralocorticoid.²³⁻²⁶ This supposition is supported by the observation that a number of such cases have low levels of plasma potassium²⁴ and an increase in total exchangeable sodium.²⁵ Crane and colleagues²⁷ treated 43 patients of this type with high doses of spironolactone and achieved a marked reduction in blood pressure in 77 per cent within three weeks whereas only 38 per cent of patients otherwise similar but with normal levels of renin showed such blood pressure reduction. This and other reports²⁷ may be regarded as circumstantial evidence in favor of excess of a circulating mineralocorticoid in some of the low renin group. In the series of Crane and associates² and Carey and co-workers²⁷ measurements of 18 hydroxy DOC and DOC were not reported and it is possible that these steroids may have been present in excess in at least some cases.

In this unit we have studied 10 cases of hypertension in whom the plasma renin concentration fell below the normal range on at least one occasion and in whom aldosterone and DOC excess were not

found. In 8 of the 10 a good hypotensive response similar to that seen in the majority of cases of primary aldosteronism was achieved. By contrast of 8 patients with essential hypertension and plasma renin concentration persistently within the normal range only one achieved a fall in diastolic pressure to below 95 mm Hg.

It appears unlikely however that all cases of hyporeninemic hypertension are associated with mineralocorticoid excess. Schalekamp and colleagues²⁸ studied 50 cases of essential hypertension excluding those with hypokalemia and found no increase in plasma and extracellular fluid volumes in the patients with low plasma renin in comparison with patients with normal levels. They did observe however that suppression of plasma renin increased in frequency with advancing years and this led to the suggestion that patients with essential hypertension and normal levels of renin may develop renin suppression as they grow older. It may well be that the 23 per cent of patients in the series by Crane and colleagues²⁷ who had low levels of plasma renin but who did not respond to spironolactone were of this type.

Spirolactone in essential hypertension

Spirolactone does have a significant albeit small hypotensive action in some cases of essential hypertension the effect being comparable only with that of thiazide diuretics.^{11, 12} The dosage of spironolactone employed in most series has generally however been smaller (150 to 200 mg per day) than that given in syndromes of definite mineralocorticoid excess (see above). The potency of large doses of spironolactone in essential hypertension remains to be fully assessed.¹⁰² There have been few clinical trials but Crane and colleagues²⁷ and Carey and associates²⁷ demonstrated a much less impressive antihypertensive effect in patients with normal renin than in those presenting with low renin hypertension.

On present evidence both spironolactone and thiazides while possibly effective alone in mild to moderate hypertension require the addition of more potent agents in the treatment of severe hypertension.¹⁰⁴ Thiazides and spironolactone when used in combination may show some potentiation

tion of renin and angiotensin II with aggravation of the hypertension. It has been demonstrated in patients of this type that renin and angiotensin II levels are disproportionately elevated in relation to the normal or only slightly reduced exchangeable sodium^{31, 32} and thus circulating angiotensin might well be sufficient to have a direct pressor effect. It is suggested that the grossly diseased kidneys provide an appropriate signal to renal release in relation to total exchangeable sodium, and that this signal is intensified when salt and water are removed at dialysis. Bilateral nephrectomy is followed by a steep fall in circulating renin and angiotensin II to very low levels and usually prompt control of the arterial pressure.^{33, 34}

SPIRONOLACTONE IN SECONDARY HYPERALDOSTERONISM Although it may be of value in correcting hypokalemia, spironolactone therapy has been generally found ineffective in controlling the hypertension in syndromes of secondary hyperaldosteronism.^{1, 22, 35, 36, 37} This is hardly surprising as aldosterone excess does not appear to be the principal cause of the hypertension. In contrast to primary hyperaldosteronism in which total exchangeable sodium is elevated and can be corrected by spironolactone therapy, in secondary hyperaldosteronism total exchangeable sodium is normal or reduced and spironolactone treatment serves only to lower total exchangeable sodium further and therefore usually to intensify the basic abnormalities. Thus while a slight and sometimes helpful hypotensive effect can be achieved, other forms of therapy need to be employed to achieve a full therapeutic effect.

Pregnancy oral contraceptive therapy

The renin/angiotensin/aldosterone system appears not to be causally related to the hypertension of pregnancy.^{44, 45} In normal pregnancy renin, renin substrate, angiotensin II, and aldosterone increase markedly, whereas in pre-eclamptic toxemia, all four components are significantly lower than in normal pregnancy, although (apart from angiotensin II) often remaining above the non-pregnant range.⁴⁶ Diuretics are frequently used in the treatment of this condition when edema is present and spironolactone has been found to be as effective as benzothiadiazines.⁴⁷ However,

the rational basis for using an aldosterone antagonist in this situation is uncertain and such therapy cannot on present evidence be recommended.

The hypertension produced in some women taking oral contraceptives does not appear to be consistently associated with aldosterone excess^{48, 49} and is best treated by stopping the tablets.

Spironolactone in syndromes of mineralocorticoid excess other than aldosterone

Genest and Nowaczynski⁵⁰ and Melby and associates⁵¹ have reported increased secretion of 18-hydroxydeoxycorticosterone in some patients with hypertension, suppressed plasma renin activity, and normal or reduced aldosterone secretion. The hypertension in some of these was reported to have been corrected by administration of spironolactone in high dosage.⁵²

It has been shown in rats⁵³ and in man⁵⁴ that the spiroclactones block the salt-retaining actions of deoxycorticosterone acetate (DOCA) as well as of aldosterone. Since DOCA is rapidly hydrolyzed to deoxycorticosterone (DOC) *in vivo*,⁵⁵ spironolactone might be expected to be effective in clinical hypertension associated with DOC excess.

Brown and colleagues⁵⁶ found that of 21 patients with hypertension associated with suppression of plasma renin, 6 had consistently elevated plasma concentration of DOC although plasma aldosterone was normal or was 11-hydroxycorticosteroids and their response to synthetic corticotrophin. The mean plasma potassium in this group of patients was low. Five of the 6 patients were treated with high doses of spironolactone and in all cases there was a significant hypotensive response which was marked in three. Hypokalemia was in all instances corrected during therapy. These observations confirmed that some cases of low renin hypertension could indeed have an excess of a previously unidentified mineralocorticoid.

In Cushing's syndrome specific therapy to correct the steroid excess is indicated⁵⁷ and spironolactone has not as far as we are aware been generally advocated in the treatment of the hypertension and electrolyte abnormalities. We have in one patient used it successfully to correct hypokalemia. Caution is necessary as spironolactone will interfere with the measurement of 11-hy

- 9 Brier L, Sommers S C, Krikoff L, Newton M A and Liragh J H Pseudo-primary aldosteronism: An entity distinct from true primary aldosteronism *Circ Res* 25 and 26 (Suppl 1) 203 1970
- 10 Bighieri E G, Schambelan M, Slaton P E and Stockigt J R The intercurrent hypertension of primary aldosteronism *Circ Res* 26 and 27 (Suppl 1) 195 1970
- 11 Brown J J, Dave D A, Ferris J B, Fraser R, Haywood E, Lever A F and Robertson J I S Comparison of surgery and prolonged spironolactone therapy in patients with hypertension, aldosterone excess and low plasma renin *Br Med J* 2 729 1972
- 12 Bighieri E C, Stockigt J R and Schambelan M Adrenal mineralocorticoids causing hypertension *Am J Med* 52 623 1972
- 13 Foye L A and Feichtmeir T A Adrenocortical carcinoma producing solely mineralocorticoid effect *Am J Med* 19 666 1955
- 14 Brooks H V, McSwiney H R, Trunty F T G and Wood F J A Potassium deficiency of renal and adrenal origin *Am J Med* 23 391 1957
- 15 Zimmerman B, Moran W H, Rosenberg J C, Kennedy B J and Frey R J Physiologic and surgical problems in the management of primary aldosteronism *Ann Surg* 150 653 1959
- 16 Conn J W, Knopf I F and Nebit R M Clinical characteristics of primary aldosteronism from an analysis of 145 cases *Am J Surg* 107:139 1964
- 17 Cone M G, Harris J J and Herber R Primary aldosteronism due to an adrenal carcinoma *Ann Intern Med* 63 494 1965
- 18 Standarter S I, Gonzalez A and Suarez J A A case of probable mineralocorticoid excess without hypercortisolism due to carcinoma of the adrenal cortex *J Clin Endocrinol Metab* 25 1429 1965
- 19 Neville A M and Symington T Pathology of primary aldosteronism *Cancer* 19:1854 1966
- 20 Harrison J H In discussion of Slen W, Bighieri E G, Slaton P et al Management of primary aldosteronism: evaluation of potassium and sodium balance, technique of adrenal ectomy and operative results in 24 cases *Ann Surg* 164 600 1966
- 21 Bighieri E G, Slaton P E, Schambelan M and Kronfield J J Hypermineralocorticoidism *Am J Med* 42 110 1968
- 22 Brock H V, Felix Davis D, Lee M R and Robertson P W Hyperaldosteronism from adrenal carcinoma *Br Med J* 1 220 1972
- 23 Brown J J, Ferris J B, Fraser R, Lever A F and Robertson J I S Aldosterone excess and adrenal carcinoma *Br Med J* 3 686 1972
- 24 Filipceki S, Feltynowski T, Poplanska W, Lapinska K, Krus S, Wocial B and Januszewicz W Carcinoma of the adrenal cortex with hyperaldosteronism *J Clin Endocrinol Metab* 32 225 1972
- 25 Sutherland D J A, Ruse J I and Laidlaw J C Hypertension increased aldosterone secretion and low plasma renin activity relieved by dexamethasone *Can Med Assoc J* 95:1109 1966
- 26 New M I and Peterson R F A new form of congenital adrenal hyperplasia *J Clin Endocrinol Metab* 27 100 1967
- 27 Mura K, Demura H, Sato F, Sasano Y and Shimizu N Acute of glucocorticoid responsive hyperaldosteronism *J Clin Endocrinol Metab* 28:1807 1968
- 28 Salt I S, Stiefel M, Ruse J I and Laidlaw J C Non tumorous primary aldosteronism: Type relieved by Glucocorticoid (Glucocorticoid Remediable Aldosteronism) *Can Med Assoc J* 101:1 1969
- 29 Barricough M A Sodium and water depletion with acute malignant hypertension *Am J Med* 40:265 1966
- 30 Brown J J, Davies D L, Lever A F and Robertson J I S Plasma renin concentration in human hypertension III Renin in relation to complications of hypertension *Br Med J* 1:50 1966
- 31 Barricough M A, Bacchus B, Brown J J, Davies D L, Lever A F and Robertson J I S Plasma renin and aldosterone secretion in hypertensive patients with renal or renal artery lesions *Lancet* 2:1310 1965
- 32 Brown J J, Lever A F and Robertson J I S Renin and angiotensin in health and disease *Schweiz Med Wochenschr* 97 1679 1967
- 33 Robertson J I S, Brown J J, Dusterdieck G O, Ferris J B, Fraser R and Lever A F Hypertension with aldosterone excess *Anglo Ger Med Rev* 6 55 1972
- 34 Brown J J, Curtis J R, Lever A F, Robertson J I S, de Wardener H H and Wing A J Plasma renin concentration and the control of blood pressure in patients on maintenance haemodialysis *Nephron* 6:329 1969
- 35 Brown J J, Dusterdieck G O, Fraser R, Lever A F, Robertson J I S, Tree M and Weir R J Hypertension and chronic renal failure *Br Med Bull* 27 128 1971
- 36 Robertson P W, Kildjian A, Harding L K, Walters G, Lee M R and Robb-Smith A H T Hypertension due to a renin secreting tumour *Am J Med* 43 963 1967
- 37 Mitchell J D, Baxter T J, Blair West J R and McCredie D A Renin levels in nephroblastoma (Wilms tumour) *Arch Dis Child* 45:376 1970
- 38 Lee M R Renin secreting kidney tumours: A rare but remediable cause of serious hypertension *Lancet* 2 254 1971
- 39 Schambelan M and Bighieri E G Regulation and significance of hyperreninemia from renin secreting tumour *Clin Res* 10 439 1972
- 40 Conn J W, Cohen E L, Lucas C P, McDonald W J, Mayor G H, Blough W M, Eveland W C, Bookstein J J and Lapidus J Primary reninism: hypertension, hyperreninemia and secondary aldosteronism due to renin producing juxtaglomerular cell tumours

of effect and there may be an additional benefit in that the spironolactone will correct hypokalemia induced by thiazides.⁹⁸

There is some evidence that in essential hypertension, the salt and water diuresis induced by thiazides is short lived whilst the antihypertensive action persists.¹⁰¹ After prolonged treatment plasma and extracellular fluid volumes^{105, 106} and cardiac output¹⁰⁷ have all been described as returning to pretreatment levels. More recently however⁴³ it has been reported that plasma volume contraction persists although this may not entirely account for the hypotensive action. It has been postulated that the thiazides may exert some of their hypotensive action by a peripheral vascular mechanism independent of the diuretic effect and there is evidence that spironolactone shares such a mechanism¹⁰⁸ with thiazides. This may explain why in essential hypertension where there is no demonstrable salt or water excess spironolactone is only partially effective, in contrast to the usually marked response seen in low renin hyperaldosteronism. If a patient with apparently essential hypertension develops a very marked reduction in blood pressure while taking spironolactone the diagnosis of mineralocorticoid hypertension should be considered.

Mode of action

As spironolactone specifically blocks the receptor sites of aldosterone, it is to be expected that this agent would be most effective where excess of aldosterone is the primary cause of the hypertension. However spironolactone has been shown to be equally effective in some syndromes of mineralocorticoid excess where the aldosterone concentration is normal but in these syndromes total exchangeable sodium is increased also.

Where the total exchangeable sodium is not elevated a less powerful hypotensive effect is likely although it is perhaps not surprising that some striking successes may occasionally be achieved. Early reports of the use of amiloride a potassium conserving diuretic which acts independently of the presence or absence of aldosterone indicate that it may produce similar correction of hypertension plasma electrolytes, and body spaces to that achieved by spironolactone, both in cases of aldo-

sterone secreting adenoma and in non aldosterone mineralocorticoid excess.^{109, 110} Whether such non specific potassium-conserving natriuretic agents¹¹¹ will be as effective as spironolactone remains to be determined.

Conclusions

While there is increasing interest in the use of spironolactone as a diagnostic and therapeutic agent in different forms of hypertension the multiple side effects although not usually severe require that high dose therapy should not be embarked upon lightly. Lower doses however may be a very effective addition to other hypotensive agents and probably are associated with less unpleasant symptoms. Extreme caution is necessary in patients with renal impairment since uremia may be aggravated and dangerous hyperkalemia provoked.

Dr D G Beever is a research fellow supported by G D Searle & Co Ltd High Wycombe England.

REFERENCES

- 1 Brown J J, Fraser R, Lever A F and Robertson J I S. Aldosterone physiological and pathophysiological variations in man. *Clinics Endocrinol Metabol* W B Saunders London 1972 19:2.
- 2 Conn J W. Primary aldosteronism: a new clinical syndrome. *J Lab Clin Med* 44:6 1955.
- 3 Brown J J, Davies D I, Lever A F and Robertson J I S. Plasma renin in a case of Conn's syndrome with bilateral lesions: use of spironolactone in treatment. *Br Med J* 2 1636 1964.
- 4 Kirkendall W M, Litz A and Armstrong M E. Hypokalemia in the diagnosis of hypertension. *Dis Chest* 45 337 1964.
- 5 Conn J W, Cohen L I and Rovner D R. Suppression of plasma renin activity in primary aldosteronism. *JAMA* 190 213 1964.
- 6 Katz F H. Primary aldosteronism with suppressed plasma renin activity due to bilateral nodular adrenocortical hyperplasia. *Ann Intern Med* 67 1035 1967.
- 7 Brown J J, Chinn R H, Davies D I, Dusterdieck C O, Fraser R, Lever A F, Robertson J I S and Tree M. Plasma electrolytes, renin and aldosterone in the diagnosis of primary hyperaldosteronism. *Lancet* 2 55 1968.
- 8 Distler A, Barth C, Roscher S, Vecsei P, Dhoni G and Wolf H J. Hochdruck und Aldosteronismus bei solitären Adenomen und bei nodulärer Hyperplasie der Nebennieren. *Deutsche Klin Wochenschr* 47 688 1969.

9. Brier L, Sommers S C, Kriskoff L, Newton M A and Liss A J H Pseudo-primary aldosteronism: An entity distinct from true primary aldosteronism. *Circ Res* 28 and 26 (Suppl 1) 203 1970
10. Biglieri E G, Shambelman M, Slaton I F and Stockigt J R The intercurrent hypertension of primary aldosteronism. *Circ Res* 26 and 27 (Suppl 1) 195 1970
11. Brown J J, Davies D I, Ferris J B, Fraser R, Haywood E, Lever A F and Robertson J I S Comparison of surgery and prolonged spironolactone therapy in patients with hypertension, aldosterone excess and low plasma renin. *Br Med J* 2 729 1972
12. Biglieri E G, Stockigt J R and Schambelman M Adrenal mineralocorticoids causing hypertension. *Am J Med* 52 673 1972
13. Foye L V and Feichtmeir T V Adrenocortical carcinoma producing solely mineralocorticoid effect. *Am J Med* 19 966 1955
14. Brooks R V, McSwiney R R, Prunty F T G and Wood F J V Potassium deficiency of renal and adrenal origin. *Am J Med* 23 391 1957
15. Zimmerman B, Moran W H, Rosenberg J C, Kennedy B J and Frey R J Physiological and surgical problems in the management of primary aldosteronism. *Ann Surg* 150 653 1959
16. Conn J W, Knopf R F and Nesbitt R M Clinical characteristics of primary aldosteronism from an analysis of 145 cases. *Am J Surg* 107 1139 1964
17. Crane M G, Harris J J and Herber R Primary aldosteronism due to an adrenal carcinoma. *Ann Intern Med* 63 494 1965
18. Standinter S R, Gonzalez A and Suarez J A Case of probable mineralocorticoid excess without hypercortisolism due to carcinoma of the adrenal cortex. *J Clin Endocrinol Metab* 25 1429 1965
19. Neville A M and Symington T Pathology of primary aldosteronism. *Cancer* 19 1854 1966
20. Harrison J H In discussion of Silen W, Biglieri E G, Slaton P et al Management of primary aldosteronism: an evaluation of potassium and sodium balance technique of adrenal ectomy and operative results in 24 cases. *Ann Surg* 164 600 1966
21. Biglieri E G, Slaton P E, Schambelman M and Kronfield J J Hypermineralocorticoidism. *Am J Med* 42 170 1968
22. Brook R V, Felix Davis D, Lee M R and Robertson P W Hyperaldosteronism from adrenal carcinoma. *Br Med J* 1 220 1972
23. Brown J J, Ferris J B, Fraser R, Lever A F and Robertson J I S Aldosterone excess and adrenal carcinoma. *Br Med J* 3 686 1972
24. Filipeck S, Feltyson L T, Poplanska W, Lapinska H, Krus S, Wocial B and Januszewicz W Carcinoma of the adrenal cortex with hyperaldosteronism. *J Clin Endocrinol Metab* 33 275 1972
25. Sutherland D J A, Ruse J I and Laidlaw J C Hypertension increased aldosterone secretion and low plasma renin activity relieved by dexamethasone. *Can Med Assoc J* 95 1109 1966
26. New M I and Peterson R F A new form of congenital adrenal hyperplasia. *J Clin Endocrinol Metab* 27 1300 1967
27. Miura K, Demura H, Sato F, Sasano Y and Shimizu Y A case of glucocorticoid responsive hyperaldosteronism. *J Clin Endocrinol Metab* 28 1807 1968
28. Salt I S, Stiefel M, Ruse J L and Laidlaw J C Non tumorous primary aldosteronism: I Type relieved by Glucocorticoid (Glucocorticoid Remediable Aldosteronism). *Can Med Assoc J* 101 11 1969
29. Barracough M A Sodium and water depletion with acute malignant hypertension. *Am J Med* 40 265 1966
30. Brown J J, Davies D I, Lever A F and Robertson J I S Plasma renin concentration in human hypertension III Renin in relation to complications of hypertension. *Br Med J* 1 505 1966
31. Barracough M A, Bacchus B, Brown J J, Davies D I, Lever A F and Robertson J I S Plasma renin and aldosterone secretion in hypertensive patients with renal or renal artery lesions. *Lancet* 2 1310 1965
32. Brown J J, Lever A F and Robertson J I S Renin and angiotensin in health and disease. *Schweiz Med Wochenschr* 97 1679 1967
33. Robertson J I S, Brown J J, Dusterdieck G O, Ferris J B, Fraser R and Lever A F Hypertension with aldosterone excess. *Anglo Ger Med Rev* 6 55 1972
34. Brown J J, Curtis J R, Lever A F, Robertson J I S de Wardener H E and Wing A J Plasma renin concentration and the control of blood pressure in patients on maintenance haemodialysis. *Nephron* 6 329 1969
35. Brown J J, Dusterdieck G O, Fraser R, Lever A F, Robertson J I S, Tree M and Weir M J Hypertension and chronic renal failure. *Br Med Bull* 27 128 1971
36. Robertson P W, Klidjian A, Harding L K, Walters G, Lee M R and Robb-Smith A H T Hypertension due to a renin secreting tumour. *Am J Med* 43 963 1967
37. Mitchell J H, Baxter T J, Blair West J R and McCredie D A Renin levels in nephroblastoma (Wilms tumour). *Arch Dis Child* 45 376 1970
38. Lee M R Renin secreting kidney tumours: A rare but remediable cause of serious hypertension. *Lancet* 2 254 1971
39. Shambelman M and Biglieri E G Regulation and significance of hyperreninemia from renin secreting tumour. *Clin Res* 20 439 1972
40. Conn J W, Cohen E L, Lucas C P, McDonald W J, Mayor G H, Blough W M, Eveland W C, Bookstein J S and Lapidus J Primary reninism, hypertension, hyperreninemia and secondary aldosteronism due to renin producing juxtaglomerular cell tumours

- Arch Intern Med 130:682 1972
- 41 Conn J W and Conn I S Primary aldosteronism versus hypertensive disease with secondary aldosteronism Recent Progr Horm Res 17:389 1961
 - 42 Brown J J Davies D L Lever A I and Robertson J I S Plasma renin concentration in human hypertension I Relationship between renin sodium and potassium Br Med J 2:144 1965
 - 43 Bourgoignie J J Catanzaro P J and Perry H M Renin-angiotensin-aldosterone system during chronic thiazide therapy for benign hypertension Circulation 37:27 1968
 - 44 Catt K J Crun L Zimmert P / Best J H Crun M D and Coughlin J P Angiotensin II blood levels in human hypertension Lancet 1:459 1971
 - 45 Farizi R C Dustan H P and Frohlich E D Long term thiazide therapy in essential hypertension: evidence for persistent alteration in plasma volume and renin activity Circulation 41:709 1970
 - 46 Weir R J Puntin D B Brown J J Fraser R Lever A I Robertson J I S and Young J A serial study in pregnancy of the plasma concentration of renin, corticosteroids, electrolytes and proteins and of haematocrit and plasma volume J Obstet Gynaecol Br Commun 78:590 1971
 - 47 Robertson J I S Weir R J Dusterdieck G O Fraser R and Tree M Renin, angiotensin and aldosterone in human pregnancy and the menstrual cycle Scott Med J 16:183 1971
 - 48 Weir R J Brown J J Fraser R Kruszewski A Lever A I McIlwaine G Morton J J Robertson J I S and Tree M Plasma renin, renin substrate, angiotensin II and aldosterone in hypertensive disease of pregnancy Lancet 1:291 1973
 - 49 Newton M A Sealey J I Iedingham J G G and Farrah J H High blood pressure and oral contraceptives: Changes in plasma renin and renin substrate and in aldosterone excretion Am J Obst Gynecol 101:1037 1968
 - 50 Weinberger M H Collins R D Dowdy A J Nokes G W and Luetscher J A Hypertension induced by oral contraceptives: continuing, estrogen and gestagen effects on plasma renin activity and aldosterone excretion Ann Intern Med 71:891 1969
 - 51 Weir R J Briggs I Mick A Faylor I Browning J Nussmith I and Wilson L Blood pressure in women after one year of oral contraception Lancet 1:467 1971
 - 52 Weir R J Free M Fraser R Chinn H H Davie D I Dusterdieck G O Robertson J I S Horne C H W and Mallinson A C The effect of combined oestrogen-progestogen oral contraceptives and of their separate components on plasma levels of renin, renin substrate, angiotensin and aldosterone and on blood pressure Proceedings of the Third International Congress on Hormonal Steroids Excerpta Medica International Congress Series No 219 1971 p 929
 - 53 Weir R J Tree M and McElwee G Changes in blood pressure and in plasma renin substrate and angiotensin II concentration in women taking contraceptive steroids Proceedings of the Fourth International Congress of Endocrinology Washington D C 1972 (In press 1973)
 - 54 Genest J and Nowaczynski W Aldosterone and electrolyte balance in human hypertension J R Coll Physicians Lond 5:77 1970
 - 55 Melby J C Dile S I and Wilson T E 18-Hydroxy-deoxycorticosterone in human hypertension Circ Res 28 and 29 (Suppl 2) 143 1971
 - 56 Brown J J Ferriss J B Fraser R Lever A I Love D R Robertson J I S and Wilson A Apparently isolated excess of deoxycorticosterone in hypertension: a variant of the mineralocorticoid excess syndrome Lancet 2:243 1972
 - 57 Goldsmith O Solomon D H and Horton R H Hypogonadism and mineralocorticoid excess N Engl J Med 277:673 1967
 - 58 Bihleri E G Herron M A and Brut W 17-hydroxylation deficiency in man J Clin Invest 45:1946 1966
 - 59 Baulieu E F Leillon I and Migeon C J Adrenogenital syndrome in Eisenstein A B ed The Adrenal Cortex London 1967 J & A Churchill Ltd p 353
 - 60 Liddle G W Bledsoe T and Coppage W S A familial renal disorder simulating primary aldosteronism but with negligible aldosterone secretion in Baulieu E F and Kober P editors Aldosterone Oxford 1964 Blackwell Scientific Publications p 351
 - 61 Fraser R James V H I London J Kurt W S Lawson A Giles C A and McKay A M Clinical and biochemical studies of a patient with a corticosterone secreting, adrenocortical tumour Lancet 2:1116 1968
 - 62 Borst J C G Blonhert G Molhuysen J A Gerbrandy J Turner K P and De Vries I A De uitcheidung van water en elektrolyten gedurende het elmaral in onder invloed van succus liquoritiae Acta Clin Belg 14:105 1950
 - 63 Molhuysen J A Gerbrandy J De Vries I A De Jong J C Ienstra J B Turner K P and Borst J C G A liquorice extract with deoxycortone-like action Lancet 2:381 1950
 - 64 Salassa R M Mattox V R and Loseveit J W Inhibition of the mineralocorticoid activity of liquorice by spironolactone J Clin Endocrinol Metab 22:1156 1962
 - 65 Conn J W Royner D H and Cohen I L Liquorice-induced pseudoaldosteronism JAMA 200:80 1968
 - 66 Holmes A M Marriott P K Young J and Irenite I Pseudoaldosteronism induced by habitual ingestion of liquorice J Clin Med J 46:625 1970
 - 67 Doll R Hill I D and Hutton C I Treatment of gastric ulcer with carbinoxalone sodium and oestrogens Gut 6:19 1965
 - 68 Luetscher J A and Lieberman A H Aldosterone Arch Intern Med 123:1958 1958

- 69 Wood J W Liddle G W Stant E G Michelakis A M and Bril A B Effect of an adrenal inhibitor in hypertensive patients with suppressed renin Arch Intern Med 123:366 1969
- 70 Spark R F and Melby J C Hypertension and low plasma renin activity presumptive evidence of mineralocorticoid excess Ann Intern Med 5 831 1971
- 71 Ledingham J G G Bull M B and Laragh J H The meaning of aldosterone in hypertensive disease Circ Res 30 and 31 (Suppl 2) 177 1967
- 72 Crane M C Harris J J and Johns V J Hyporeninaemic hypertension Am J Med 43:457 1972
- 73 Schalekamp M A D H Kraus V H Holsters G Schalekamp M P A and Birkenhager W H Renin suppression in hypertension in relation to body fluid volumes patterns of sodium excretion and renal haemodynamics Clin Sci (In press 1973)
- 74 Brown J J Davies D L Lever A F Peart W M and Robertson J I S Plasma concentration of renin in a patient with Conn's syndrome and fibroid lesions of the renal arterioles the effect of treatment with spironolactone J Endocrinol 33 279 1965
- 75 Brown J J Fraser R Lever A F and Robertson J I S Hypertension with aldosterone excess Br Med J 2:391 1972
- 76 Fraser R Brown J J Chinn R H Lever A F and Robertson J I S The control of aldosterone secretion and its relationship to the diagnosis of hyperaldosteronism Scott Med J 14:420 1969
- 77 Brown J J Fraser R Lever A F and Robertson J I S Hypertension a review of selected topics Abstracts of World Medicine 40:349 1971
- 78 Brown J J Dusterdieck G O Ferriss J B Fraser R Lever A F and Robertson J I S The renin-angiotensin system in hypertension Seventh Symposium on Advanced Medicine Boucher I A editor London 1971 Pitman Medical Publishing Co Ltd p 265
- 79 Brown J J Davies D L Lever A F and Robertson J I S Plasma renin concentration in human hypertension II Renin in relation to aetiology Br Med J 2:1215 1965
- 80 Spark R F and Melby J C Aldosteronism in hypertension The spironolactone response test Ann Intern Med 69 685 1968
- 81 Ferriss J B Brown J J Fraser R Kay A W Neville A M O Murchieartaigh I G Robertson J I S Symington T and Lever A F Hypertension with aldosterone excess and low plasma renin pre-operative distinction between patients with and without adrenocortical tumour Lancet 2 995 1970
- 82 Crane M C and Harris J J Effect of spironolactone in hypertensive patients Am J Med Sci 260:311 1970
- 83 Mattingly M A simple fluorimetric method for estimation of 11 hydroxycorticoids in human plasma J Clin Pathol 15:374 1962
- 84 Brown J J Ferriss J B Fraser R Lever A F and Robertson J I S Spironolactone in the treatment of hypertension with aldosterone excess in Wilson G M editor Medical uses of spironolactone Amsterdam 1971 Excerpta Medica p 27
- 85 Atchison J Brown J J Ferriss J B Fraser R Kay A W Lever A F Neville A M Symington T and Robertson J I S Quadratic analysis in the preoperative distinction between patients with and without adrenocortical tumours in hypertension with aldosterone excess and low plasma renin Am Heart J 82 660 1971
- 86 Pickering G W In High blood pressure 2nd ed London 1968 J & A Churchill Ltd p 496
- 87 Medina A Bell P R F Briggs J D Brown J J Fine A Lever A F Morton J J Paton A M Robertson J I S Tree M Waite M A Weir R J and Winchester J F Changes in blood pressure renin and angiotensin following bilateral nephrectomy in patients with chronic renal failure Br Med J 4 694 1972
- 88 Waite M A Circulating levels of angiotensin I and II in various hypertensive states Scand J Clin Lab Invest 2 (Suppl 29 section 7) 3 1972
- 89 Morton J J and Waite M A Radioimmunoassay of angiotensin I and II concurrent measurements of circulating levels in physiological and pathological states Eur J Clin Invest 2:297 1972
- 90 Brown J J Matthew G K and Robertson J I S The effect of angiotensin on the function of the separate kidneys in patients with unilateral renal artery stenosis Clin Sci 26 381 1964
- 91 Davies D L Beevers D G Brown J J Fraser R Ferriss J B Lever A F Medina A Morton J J and Robertson J I S Sodium and the renin-angiotensin system in patients with hypertension Proceedings of the Fourth International Congress of Endocrinology Washington 1972 (In press 1973)
- 92 Mackay E Khoo J K Adsett G and Zymanski A A clinical study of the effects of an aldosterone antagonist (Aldactone A) and a thiazide diuretic (Enduron) in pregnancy Aust N Z J Obstet Gynaecol 9:188 1969
- 93 Kagawa C M Cella J A and Van Arman C G Action of new steroids in blocking effects of aldosterone and deoxy corticosterone on salt Science 126 1015 1957
- 94 Liddle G W Sodium diuresis induced by steroidal antagonists of aldosterone Science 126 1016 1957
- 95 Fraser R and Wilson A Personal communication 1972
- 96 Berger G M and Edwards C R W Cushing's syndrome Clinics Endocrinol Metab W H Saunders London 1:451 1972
- 97 Carey R M Douglas J G and Liddle G W Spironolactone in patients with essential hypertension and suppressed plasma renin activity Clin Res 20 72 1972
- 98 Wimer B M Lubbe W F and Colton T

- Antihypertensive actions of diuretics comparative study of an aldosterone antagonist on a thiazide alone and together JAMA 204:775 1968
- 99 Cranston W I and Juel Jensen H E The effects of spironolactone and chlorthalidone on arterial pressure Lancet 1:1161 1962
 - 100 Conway J and Palmero J C The vascular effect of the thiazide diuretics Arch Intern Med 111:203 1963
 - 101 Hollander W Chobanian A V and Wilkins R W The role of diuretics in the management of hypertension Ann NY Acad Sci 88:975 1960
 - 102 Georgopoulos A J Dustan H and Page J H Spironolactone in hypertensive patients Arch Intern Med 108:389 1961
 - 103 Kellett R J Ballantyne D Malcolm M and Wapshaw J A Double-blind trial of hydrochlorothiazide and high dose spironolactone in the treatment of hypertension (In preparation 1973)
 - 104 Beavers D G Hamilton M and Harpur J E The long term treatment of hypertension with thiazide diuretics Postgrad Med J 47: 639 1971
 - 105 Wilson I M and Freis E D The relationship between plasma and extracellular fluid volume depletion and the anti hypertensive effect of chlorothiazide Circulation 20:1028 1959
 - 106 Gifford R W Mattox V R Orvis A L Sones D A and Rosevear J W Effect of thiazide diuretics on plasma volume electrolytes and excretion of aldosterone in hypertension Circulation 24:1197 1961
 - 107 Conway J and Lauwers P Hemodynamic and hypotensive effects of long term therapy with chlorothiazide Circulation 21:21 1966
 - 108 Mendlowitz M Naftchi N E Gutlow S E and Wolf R L The effect of spironolactone on digital vascular reactivity in essential hypertension AM HEART J 76:795 1968
 - 109 Kremer D Brown J J Davies D L Fraser R Lever A F and Robertson J I S Prolonged Amiloride therapy in a case of primary hyperaldosteronism with chronic peptic ulceration Br Med J (In press May 1973)
 - 110 Kremer D Beavers D G Brown J J Ferriss J B Fraser R Lever A F and Robertson J I S Spironolactone and Amiloride in the treatment of low renin hyperaldosteronism and related syndromes Clin Sci (In press May 1973)
 - 111 Antcliff A C Beavers D G Hamilton M and Harpur J E The use of amiloride hydrochloride in the correction of hypokalaemic alkalosis induced by diuretics Postgrad Med J 47:644 1971
 - 112 Leading article Primary excess and deficiency of renin Br Med J 1 627 1973
 - 113 Davies D L Schalekamp M A Beavers D G Brown J J Briggs J D Lever A F Medina A Morton J J Robertson J I S and Tree M Abnormal relation between exchangeable sodium and the renin angiotensin system in malignant hypertension and in hypertension with chronic renal failure Lancet 1 683 1973
 - 114 Morimoto S Takeda R and Murakami M Does prolonged pretreatment with large doses of spironolactone hasten a recovery from juxtaglomerular adrenal suppression in primary aldosteronism? J Clin Endocrinol 31 659 1970

Fundamentals of clinical cardiology

Peripheral arterial occlusion in patients with acute coronary heart disease

Sandor A. Friedman MD FACP*
Mahendra Pandya MD**
Ernst Greif MD FACP***
Brooklyn N Y

Relatively little is known about the epidemiology of peripheral arteriosclerosis obliterans. Although it is associated with many of the same risk factors as coronary and cerebrovascular disease accurate prevalence figures in differing populations are not available. In the past case finding in peripheral arterial disease has been hampered by lack of a precise definition of normality and in many instances reliance has been placed solely on a history of intermittent claudication. However there are now sufficient data from normal subjects to establish clinical criteria for the presence of asymptomatic arterial occlusion.^{1,2} The purpose of the present study was to apply these criteria to patients with acute coronary heart disease in order to assess the co prevalence of atherosclerosis in the extremities and the heart.

Material and methods

One hundred consecutive patients admitted to the Coronary Care Unit of the

Coney Island Hospital with acute myocardial infarction but no evidence of hypotension or significant peripheral edema were examined carefully for the presence of peripheral arterial disease. The diagnosis of acute myocardial infarction was based on characteristic symptoms in association with established electrocardiographic criteria and/or elevation of serum enzyme levels (SGOT, LDH and CPh). Every patient in the study had an abnormal electrocardiogram.

A careful history was obtained with reference to the following previous myocardial infarction symptoms compatible with congestive heart failure, angina pectoris, hypertension, diabetes mellitus, intermittent claudication, cigarette smoking.

Arterial examination in each case was performed independently by two examiners between the second and fourth hospital days. The radial, ulnar, femoral, popliteal, dorsalis pedis and posterior tibial pulses were graded as normal or

Friedman SA, Pandya M, Greif E. Peripheral arterial disease in patients with acute myocardial infarction. *Am J Med* 1973; 35: 123-128.
Reprinted with permission from the Medical Service, Coney Island Hospital, Brooklyn, N.Y. 11235.
Chapman J, et al. *Disorders of the Cardiovascular System*. New York: McGraw-Hill, 1973.
**Friedman SA, Pandya M, Greif E. *Am J Med* 1973; 35: 123-128.
***Chapman J, et al. *Disorders of the Cardiovascular System*. New York: McGraw-Hill, 1973.

Table I Summary of clinical data

| Croup | No | No of females | Mean age (yr) | History of hypertension | History of smoking |
|--|-----|---------------|---------------|-------------------------|--------------------|
| Definite peripheral arterial occlusion | 55 | 24 | 62 (41-83) | 18 (33%) | 37 (67%) |
| Normal peripheral arterial examination | 45 | 6 | 59 (41-72) | 19 (42%) | 26 (58%) |
| Total | 100 | 30 | 60 (41-83) | 37 | 63 |

absent and oscillometric readings were obtained at each ankle. With the patient lying supine systolic blood pressures were obtained by palpation of the radial pulses with a sphygmomanometer cuff around the arm and of both the dorsalis pedis and posterior tibial pulses (if present) with the cuff placed on the lower third of the calf. Systolic pressure was taken as the minimum pressure required to obliterate the pulse. Fifteen normal subjects (ages 24 to 31 years) were examined in the identical fashion.

All pulses were repeatedly examined by both observers until definite agreement could be reached. The posterior tibial pulse was sought with the patient supine and relaxed and the observer seated on the side being examined. The entire area under the medial malleolus was palpated and this maneuver was repeated with the foot externally rotated before the posterior tibial pulse was declared absent. The entire anterior surface of the foot was palpated for the dorsalis pedis artery in positions of flexion and extension.

The following findings were considered diagnostic of arterial occlusion: (1) absence of one or both posterior tibial pulses; (2) absence of radial and ulnar pulses in a wrist; and (3) a posterior tibial artery blood pressure at least 20 mm Hg lower than that of the radial arteries.

Blood sugar level at the time of hospitalization, serum cholesterol concentration, and hematocrit were also recorded for each patient.

Results

Table I summarizes the data for 55 patients with definite peripheral arterial

occlusion and for the 45 patients who did not meet the criteria for this diagnosis. Mean age was 60 years and 70 per cent were men. Of the entire series of 100 patients, 63 had been smoking for a mean of 31 years. 21 had diabetes mellitus and 34 had a history of hypertension. At least one of these 3 factors was present in 85 per cent of these patients with acute myocardial infarction. Eighty-eight per cent had one of these factors and/or definite peripheral arterial disease.

There were statistically significant differences between the sexes in this series. Twenty-four of 30 women (80 per cent) had definite peripheral arterial occlusion compared to 31 of 70 men (44 per cent) ($P < .01$). Eighty-four per cent of men and 40 per cent of women were cigarette smokers ($P < .01$). Thirty-three per cent of women had a history of diabetes mellitus as compared to 16 per cent of the men but this difference was not statistically significant (P between .05 and .10). Mean age and serum cholesterol levels were virtually identical for men and women.

A history of diabetes mellitus was present in 18 patients (29 per cent) with extremity involvement whereas 3 of 45 patients (7 per cent) were diabetic in the group without peripheral arterial occlusion ($P < .01$). Of the 21 diabetics, 4 had been treated with insulin, 12 with oral hypoglycemic agents and 5 with diet alone. No significant difference was found between the 2 groups in regard to age, history of smoking or hypertension, blood sugar or serum cholesterol level, hematocrit or mean blood pressure while in the coronary care unit.

A previous history of heart disease was obtained from 51 patients (angina pectoris

| History of diabetes | Mean serum cholesterol (mg/100 ml) | Mean blood sugar (mg/100 ml) | Hematocrit | Previous heart disease | No. with serum cholesterol over 300 mg/100 ml |
|---------------------|------------------------------------|------------------------------|------------|------------------------|---|
| 18 (33%) | 228 | 151 | 44% | 31 | 6 |
| 3 (7%) | 243 | 148 | 41% | 20 | 4 |
| 21 | 234 | 150 | 42% | 51 | 10 |

in 47 previous myocardial infarction in 31 and congestive heart failure in 14) Of the 49 patients suffering their first symptoms of heart disease 24 (49 per cent) had definite peripheral arterial occlusion. Patients with and without a history of previous heart disease were similar in all other parameters measured.

The distribution of abnormalities in patients meeting the criteria for peripheral arterial occlusion is given in Table II. In 44 cases one or both posterior tibial pulses were missing. Ten patients were diagnosed on the basis of low blood pressures in the posterior tibial arteries. Seven of these 10 also had either missing dorsalis pedis pulses or low blood pressure readings on palpation of that artery. A history of intermittent claudication was present in 18 (33 per cent) of this group. Forty-seven patients had oscillometric readings below 2 units and the other 8 patients had a normal blood pressure in the anterior tibial artery of each leg (as measured by dorsalis pedis palpation).

In addition to the 55 patients with definite arteriosclerosis of the extremities 7 patients (6 men and 1 woman) had asymmetry of their dorsalis pedis pulses. In 3 a dorsalis pedis pulse was unilaterally absent. 3 others had a 25 to 30 mm Hg pressure difference between the 2 dorsalis pedis arteries, the lower one being well below arm pressures and 1 had absence of one dorsalis pedis pulse and a low pressure in the other.

In the 15 normal subjects tibial artery blood pressure readings were never lower than those of the upper extremities. Fifty-nine of the 60 tibial arteries had pressures 10 to 25 mm Hg higher than those of the upper extremities.

Table II Findings in 55 patients with peripheral arterial occlusion

| Finding | No. |
|---|-----|
| Intermittent claudication | 18 |
| Absent posterior tibial pulse | 44 |
| Pulseless wrist | 1 |
| Low posterior tibial blood pressures (both pulses palpable) | 10 |
| Low or absent oscillometric readings | 47 |
| Femoro-popliteal arterial occlusion | 21 |

Discussion

The accessibility of the peripheral circulation to physical examination affords an opportunity to make a clinical diagnosis of asymptomatic arteriosclerosis. Previous work has shown that a missing posterior tibial or radial pulse is virtually always pathologic.^{1,2} Dorsalis pedis pulses have been reported missing in 5 to 12 per cent of normal subjects but unilaterally in only 3 to 67 per cent.^{1,2} In normal individuals intra-arterial blood pressure measurements are identical in the brachial and femoral arteries. With a sphygmomanometer cuff blood pressure is often recorded artificially higher in the lower extremities because of their wider girth but their pressures should never be lower than those of the arms.^{4,5} This was confirmed by the findings in our normal subjects. Oscillometry, although only semiquantitative, is useful for confirming the presence of peripheral arterial disease in many cases (47 of 55 in the present study). The reading decreases as perfusing pressure in the leg falls but the latter will generally be relatively unaffected if one tibial vessel remains uninvolved.⁴

Table I Summary of clinical data

| Group | No | No of females | Mean age (yr) | History of hypertension | History of smoking |
|--|-----|---------------|---------------|-------------------------|--------------------|
| Definite peripheral arterial occlusion | 55 | 24 | 62 (43-83) | 18 (33%) | 37 (67%) |
| Normal peripheral arterial examination | 45 | 6 | 59 (41-72) | 19 (42%) | 26 (58%) |
| Total | 100 | 30 | 60 (41-83) | 37 | 63 |

absent and oscillometric readings were obtained at each ankle. With the patient lying supine, systolic blood pressures were obtained by palpation of the radial pulses with a sphygmomanometer cuff around the arm and of both the dorsalis pedis and posterior tibial pulses (if present) with the cuff placed on the lower third of the calf. Systolic pressure was taken as the minimum pressure required to obliterate the pulse. Fifteen normal subjects (ages 24 to 31 years) were examined in the identical fashion.

All pulses were repeatedly examined by both observers until definite agreement could be reached. The posterior tibial pulse was sought with the patient supine and relaxed and the observer seated on the side being examined. The entire area under the medial malleolus was palpated and this maneuver was repeated with the foot externally rotated before the posterior tibial pulse was declared absent. The entire anterior surface of the foot was palpated for the dorsalis pedis artery in positions of flexion and extension.

The following findings were considered diagnostic of arterial occlusion: (1) absence of one or both posterior tibial pulses; (2) absence of radial and ulnar pulses in a wrist; and (3) a posterior tibial artery blood pressure at least 20 mm Hg lower than that of the radial arteries.

Blood sugar level at the time of hospitalization, serum cholesterol concentration, and hematocrit were also recorded for each patient.

Results

Table I summarizes the data for 55 patients with definite peripheral arterial

occlusion and for the 45 patients who did not meet the criteria for this diagnosis. Mean age was 60 years and 70 per cent were men. Of the entire series of 100 patients, 63 had been smoking for a mean of 31 years. 21 had diabetes mellitus and 34 had a history of hypertension. At least one of these 3 factors was present in 85 per cent of these patients with acute myocardial infarction. Eighty-eight per cent had one of these factors and/or definite peripheral arterial disease.

There were statistically significant differences between the sexes in this series. Twenty-four of 30 women (80 per cent) had definite peripheral arterial occlusion compared to 31 of 70 men (44 per cent) ($P < .01$). Eighty-four per cent of men and 40 per cent of women were cigarette smokers ($P < .01$). Thirty-three per cent of women had a history of diabetes mellitus as compared to 16 per cent of the men, but this difference was not statistically significant (P between .05 and .10). Mean age and serum cholesterol levels were virtually identical for men and women.

A history of diabetes mellitus was present in 18 patients (29 per cent) with extremity involvement whereas 3 of 45 patients (7 per cent) were diabetic in the group without peripheral arterial occlusion ($P < .01$). Of the 21 diabetics, 4 had been treated with insulin, 12 with oral hypoglycemic agents and 5 with diet alone. No significant difference was found between the 2 groups in regard to age, history of smoking or hypertension, blood sugar or serum cholesterol level, hematocrit or mean blood pressure while in the coronary care unit.

A previous history of heart disease was obtained from 51 patients (angina pectoris

diagnosing coronary artery disease. Since 88 per cent of the patients had peripheral arterial disease and/or a history of smoking, diabetes or hypertension, the value of these factors in evaluating a patient with atypical chest pain is obvious. Clearly, examination of the peripheral arteries is a useful screening procedure for the identification of individuals prone to coronary heart disease; they can then be studied further with a view to eliminating some of the risk factors.

Summary

One hundred consecutive patients with acute coronary heart disease were examined carefully for the presence of peripheral arterial occlusion. In addition to pulse palpation, blood pressures and oscillographic recordings were obtained in the lower extremities. Fifty-five had definite arterial occlusion and seven had findings strongly suggestive of peripheral arterial disease. Only one third of the patients with arterial occlusion gave a history of intermittent claudication. One half of the patients with no previous cardiac history already had signs of peripheral arteriosclerosis. The data suggest that examination of the peripheral circulation is a useful screening procedure for the identification of individuals prone to coronary heart disease.

REFERENCES

1. Barnhorst D A and Barner H B. Prevalence of congenitally absent pedal pulses. *N Engl J Med* 278:764 1968.
2. Stephens G. Palpable dorsalis pedis and posterior tibial pulses. *Arch Surg* 83:662 1967.
3. Friedman S A. Prevalence of palpable wrist pulses. *Br Heart J* 32:316 1970.
4. Strandness D E Jr and Bell J W. Peripheral vascular disease: diagnosis and objective evaluation using a mercury strain gauge. *Ann Surg (Suppl)* 134 1965.
5. Spittell J A Jr and Hines C A Jr. A comparison of arm and thigh blood pressures in patients with abdominal aortic aneurysms. *Angiology* 11:1 1960.
6. Kannel W B, Dawber T R, Friedman G D, Glennon W E and McNamara P M. Risk factors in coronary heart disease: an evaluation of several serum lipids as predictors of coronary heart disease. The Framingham Study. *Ann Intern Med* 61:888 1964.
7. Doyle J T, Dawber T R, Kannel W B, Kinch S H and Kahn H A. The relationship of cigarette smoking in coronary heart disease: the second report of the combined experience of the Albany NY and Framingham Mass studies. *JAMA* 190:886 1964.
8. Juergens J L, Barker N W and Hines E A Jr. Arteriosclerosis obliterans: review of 520 cases with special reference to pathogenic and prognostic factors. *Circulation* 21:188 1960.
9. Coronary heart disease in adults—United States 1960-1962. U S Department of Health Education and Welfare. National Center for Health Statistics.
10. Blood pressure of adults by age and sex—United States 1960-1962. U S Department of Health Education and Welfare. National Center for Health Statistics.
11. Mahmoud A H. A population sample study of peripheral occlusive arterial disease. Thesis. University of Michigan. Ann Arbor 1964.
12. Kannel W B, Skinner J J Jr, Schwartz M J and Shusteff D. Intermittent claudication: incidence in the Framingham Study. *Circulation* 41:875 1970.
13. McDonald L. Ischemic heart disease and peripheral occlusive arterial disease. *Br Heart J* 15:101 1953.
14. Boyd A M. The natural course of arteriosclerosis of the lower extremities. *Angiology* 11:10 1960.
15. Dry T J and Hines C A Jr. The role of diabetes in the development of degenerative vascular disease with special reference to the incidence of retinitis and peripheral neuritis. *Ann Intern Med* 14:1893 1941.
16. Bell E T. Incidence of gangrene of extremities in nondiabetic and in diabetic persons. *Arch Pathol* 49:469 1950.

Since the histopathology of atherosclerosis in the femoral and popliteal arteries is indistinguishable from coronary artery disease, it is not surprising that peripheral arterial and coronary heart diseases are intimately associated. Although epidemiologic studies on incidence and prevalence have identified risk factors common to both, such as hyperlipidemia,⁶ diabetes mellitus, and cigarette smoking,^{7,8} there is a dearth of data concerning the frequency of peripheral arterial disease. The National Center for Health Statistics has published age-related estimates of the prevalence of coronary disease⁹ and hypertension¹⁰ but offers no comparable statistics for peripheral arterial occlusion.

Most of the available information is confined to prevalence and incidence of intermittent claudication. Mahmoud¹¹ estimated that 6 per cent of men and 3 per cent of women between the ages of 55 and 74 have claudication. Kannel and co-workers¹² in following the Framingham Population Study Group noted that the risk of developing claudication was 8 to 10 times as high in people with coronary heart disease as those with no heart disease. Men with claudication developed coronary disease 3 times as frequently as those without this symptom and women 9 times as frequently. McDonald¹³ found a common coexistence of angina pectoris and intermittent claudication in 137 patients. Boyd¹⁴ noted nonfatal myocardial infarction within 5 years of the onset of claudication in about 20 per cent of 1,440 patients.

Intermittent claudication is likely to identify only a fraction of all patients with peripheral arterial disease since this symptom is usually not present unless there is involvement of the larger arteries. Furthermore, claudication will occur in a mild case of peripheral arterial disease only if the patient is in the habit of walking a considerable distance without stopping. The presence of intermittent claudication in only one third of our subjects with definite peripheral arterial occlusion is striking proof of the insensitivity of this symptom as an epidemiologic tool.

The sex distribution of peripheral arterial occlusion in patients with coronary heart disease is difficult to explain. A significantly higher propensity for generalized arterial

disease in females found in this series and the Framingham Study¹⁵ could not be explained by any difference in mean age, serum cholesterol level, smoking history, or prevalence of diabetes mellitus. Perhaps it is a reflection of a fundamental difference between men and women in the nature of atherosclerosis. Women who develop coronary heart disease in middle age may represent a group with a strong genetic predisposition to atherosclerosis and therefore might tend to have wider involvement of the arterial system, whereas this disease process in men may be more or less a natural result of the aging process.

The much higher prevalence of diabetes in patients with extremity involvement is consistent with the known susceptibility of diabetics to peripheral arteriosclerosis obliterans. The latter is estimated to be 11 to 40 times as frequent in the diabetic as in the non-diabetic population.^{13,16}

The results of the present study suggest that the majority of patients who are hospitalized with acute coronary heart disease already have clinically detectable peripheral arterial disease. In addition to the 51 patients with definite criteria for this diagnosis, 7 other patients had marked discrepancies between their 2 dorsalis pedis pulses—findings suggestive although not diagnostic of arterial occlusive disease.¹⁷ Forty-nine patients gave no history of angina, previous myocardial infarction, or congestive heart failure, and it is reasonable to presume that they were having their initial clinical manifestations of heart disease. Although it is possible that careful examination prior to their myocardial infarction would have revealed evidence of cardiomegaly or gallop rhythm, heart disease would probably have gone undetected in most cases by history and physical examination. On the other hand, significant atherosclerosis could have been diagnosed before cardiac symptoms occurred in about half these patients with close attention to the peripheral pulses. The incorporation of blood pressure measurements in the lower extremities appears to offer an advantage over pulse palpation alone since peripheral arterial disease was diagnosed by blood pressure readings alone in 10 patients.

The results of this survey suggest that certain historical information is helpful in

subjected to cardiac catheterization for suspected anginal syndrome. Indeed a small number of patients with documented myocardial infarctions have normal coronary arteriograms.

Evaluation of the function of the left ventricle is essential in the proper selection of candidates. Operative and late death as well as the effectiveness of surgery is closely related to the performance of the left ventricle. The size and shape of the chamber, its contractility, filling pressure and output are parameters usually measured. By means of left ventriculography the dynamic changes of the contracting ventricle can be appreciated and areas of reduced or absent contractility delineated. Associated mitral regurgitation can be assessed and ventricular volumes calculated. The ejection fraction that is the amount of the diastolic volume reduced with each beat is inversely proportional to the operative and late mortality rate of surgery. As the ejection fraction falls and other evidence of poor left ventricular function appears the risks of operation increase. Ventricular dysfunction is one of the most important determinants of patient survival. An operative mortality rate of less than 5 per cent in patients with good ventricular function can be 30 to 40 per cent in the presence of severe cardiac enlargement and failure.

Anginal syndrome. Anginal syndrome refractory to optimum medical treatment constitutes the principal indication for bypass surgery today. Refractoriness is an arbitrary term which must be applied to each particular patient. It is affected by the tolerability of his symptoms and its effects upon his work activities and total life style. Mild recurrent chest discomfort readily responsive to medications such as nitroglycerine may be tolerated by many individuals. Severe recurrent exertional pain, nocturnal and rest angina unrelieved by medical management generally is not acceptable.

The presence of right coronary, left anterior descending and left circumflex obstructions singly or in combination in a patient with unstable angina is an indication for operation. Complete or partial relief of symptoms may be expected in 80 to 90 per cent of these cases.

Unstable angina. Infarction angina

impending myocardial infarction, acute coronary insufficiency, intermediate coronary syndrome and crescendo angina are terms applied to an unstable anginal syndrome manifest by rapidly progressively severe and protracted chest pains. Early reports of poor prognosis and progression to infarction in this group have been questioned by more recent studies. Differences in patient selection and availability of observation and management in coronary care units may be factors in their low incidence of early infarction. Lack of arteriographic confirmation in most instances makes further comparisons even more difficult. When angiographic data are available the distribution and extent of stenotic lesions is similar in both chronic stable and acute unstable angina.

Most patients with acute development of angina will respond to rapid institution of bed rest and medical therapy. Those with sudden exacerbation of chronic angina and those in whom angina persists despite intensive management should be considered high risk patients and evaluated by arteriography for possible emergency bypass surgery.

Chronic congestive heart failure. Chronic heart failure is the predominant manifestation of many patients with coronary occlusive disease. Bypass surgery with and without resection of aneurysmal portions of the left ventricle has had little effect in regard to improvement in ventricular filling pressures, contractility and cardiac output. Surgical mortality rate is high and one year survival low.

Some patients with significant anginal syndrome and a moderately reduced left ventricular ejection fraction might be considered for bypass surgery despite the higher risks of operation. Individuals with little or no angina and severe impairment of left ventricular function should not be subjected to coronary bypass surgery with the expectation of meaningful clinical improvement.

Acute myocardial infarction. Documented acute myocardial infarction generally has been considered a contraindication to surgery. Established muscle necrosis will not be reversed by revascularization and the risks of operation are excessive. On the other hand supported by animal experi-

Indications for aortocoronary artery bypass surgery

Alfred J Kallman, M D
New York N Y

Aortocoronary artery bypass grafts for coronary occlusive disease clinically introduced in 1967 have reached epidemic proportions. As the current trend mushroomed in 1973 we should anticipate many more procedures than the estimated 20 000 performed in the United States last year.

There is no question of the efficacy of coronary bypass surgery for the relief of angina in selected patients. Whether future myocardial infarctions will be prevented and life prolonged is not yet clearly demonstrated. In this regard a randomized prospective study already is under way within the Veterans Administration Hospital system and a cooperative investigation by a number of large medical centers is being organized by the National Heart and Lung Institute. Hopefully these and the objective results of other operating groups will provide a meaningful therapeutic evaluation of the operation on the course of arteriosclerotic coronary occlusive disease in the future.

At the present time there is unbridled enthusiasm and acceptance of the procedure on the one hand, concern and caution on the other. The controversy has been a favorite subject for editorials in medical journals and the lay press has given it abundant coverage. Television and national

periodicals have informed the public of its potential and increased the pressures of patient demand.

Objective evaluation of the effects of surgery vary depending upon the specific technique of the procedure, selectivity of patients, effective follow up study and the natural course of the disease. There is a tremendous emotional bias in any life threatening situation and the possibility of amelioration of a destructive process by an aggressive approach even if not completely proved can be appealing to the afflicted patient and his physician. It is in this emotionally charged atmosphere that the physician must decide if his patient is or is not a candidate for coronary bypass surgery today.

Coronary arteriography with demonstration of significant proximal large vessel occlusion is paramount in the consideration for surgery. Obstruction generally is considered significant when the reduction in size of the lumen is greater than 75 per cent. Visualization of normal arterial distribution distal to the stenosis is desirable but not essential. At operation patent distal vessels not apparent angiographically may be found suitable for bypass grafting. Lack of coronary occlusive disease may be found in 15 to 27 per cent of various series of patients.

From the Department of Medicine, New York University Medical Center, New York, N. Y.

Received for publication May 21, 1973.

Reprint requests to Dr. Alfred J. Kallman, Department of Medicine, New York University Medical Center, New York, N. Y. 10016.

therapy and the presence of severe angina in most of the patients would seem to warrant consideration for early operation. This lesion is not common, occurring in less than 3 per cent of patients undergoing coronary arteriography, and experience with surgery is not extensive. Further objective evaluation is necessary, particularly since left main arterial stenosis is almost never a solitary occlusive process.

Asymptomatic patients In general asymptomatic patients despite documented coronary occlusive disease are not operated upon except under certain special circumstances. Coronary arteriography in the assessment of valvular aortic stenosis may disclose morphologically significant obstructions in patients without angina. Coronary bypass is not performed routinely, unless there is difficulty in weaning the patient off the pump-oxygenator after prosthetic replacement of the valve or there is severe compromise of the coronaries with poor ventricular perfusion and anticipated low output failure in the early postoperative period. Similar situations may arise during ventricular aneurysmectomy and if feasible revascularization using one or more grafts may be attempted.

There is inadequate information as regards the benefits of surgery in the asymptomatic patient with uncomplicated prior myocardial infarction or demonstrable limited coronary artery disease without infarction. Prediction of impending myocardial infarction or sudden death in the asymptomatic patient is not yet possible. As high risk factors are identified in the natural course of the disease and further correlations are made with coronary arteriography and left ventricular function studies, perhaps a more liberal selectivity of candidates for surgery will follow.

Risk factors Hypertension, smoking, hypercholesterolemia and hyperlipidemia have been associated with acceleration of coronary atherosclerosis. The over all yearly mortality rate of 3 to 4 per cent in patients with angina is increased to 8 per cent in the sub group with high blood pressure and advanced electrocardiographic abnormalities. Cardionegaly and congestive heart failure adversely affect the prognosis.

With coronary arteriography we can attempt correlation of survival with dis-

tribution and extent of arterial stenosis. Multiple vessel disease particularly involving the left anterior descending branch worsens the prognosis. Approximately 90 per cent of patients with single vessel disease have a two year survival, only 60 to 70 per cent with two and three vessel involvement live that long. The importance of coronary collateral circulation and progression of obstructive lesions are variables with insufficient data for meaningful evaluation at this time.

As increase in vascular stenoses and reduction in left ventricular function influence the natural history of coronary arterial disease, these same factors increase the operative risks as well. The intraoperative and postoperative mortality rate and infarction rate are higher in those patients requiring multiple bypass grafts and in those with ventricular dysfunction.

In addition to the surgical mortality rate ranging from 3 to 10 per cent, we must consider the effectiveness of operation. The patency of saphenous vein aortocoronary grafts is in the range of 80 per cent after two years. A one year patency of 97 per cent has been claimed for internal mammary artery bypass, a procedure requiring longer operating time and with an obviously limited supply of vessels for grafting.

Conclusions

Adequate statistically significant randomized data is not yet available to determine the effect of aortocoronary bypass grafting on the natural course of occlusive coronary atherosclerosis. Large prospective studies under way or in the planning stage may help define the influence of surgery upon future myocardial infarction and longevity.

There is no doubt of the relief of angina in about 80 per cent of patients undergoing operation. In the absence of left ventricular failure, operative mortality rate is low and if angina is incapacitating and unresponsive to medical management, surgery should be considered. Left main coronary lesions, unstable angina with acute unrelieved exacerbation and myocardial infarction with cardiogenic shock are high risk situations in which operation if feasible would be recommended.

Bypass surgery has been disappointing

mentation, marginal areas may be improved and the extent of necrosis may be reduced if the vascular supply is restored within three hours of occlusion.

Patients developing coronary occlusion during or shortly after coronary arteriography have been operated upon satisfactorily within hours of the event. However, this surgical experience with persons sustaining coronary occlusion in a hospital setting cannot be applied to those admitted with myocardial infarction which occurred even a few hours prior to admission. The further delay inherent in the necessary preoperative assessment by cardiac catheterization would be enough to preclude the possibility of reversibility of the developing myocardial necrosis. At the present time surgery for uncomplicated acute myocardial infarction cannot be recommended.

Continued angina during the course of acute myocardial infarction may be considered a variant of the unstable anginal syndrome. It would suggest additional areas of ischemia in jeopardy. If this is severe and is unresponsive to conservative measures the possibility of surgical intervention should be evaluated by coronary arteriography. The risk of additional infarction might outweigh that of operation in these patients with otherwise uncomplicated acute myocardial infarction.

Acute myocardial infarction with cardiogenic shock. The high mortality rate in the ninety percentile range associated with the cardiogenic shock of acute myocardial infarction has prompted aggressive treatment in several small series of patients. Mechanical circulatory assistance with an intra-aortic balloon pump has been effective in some instances but the mortality rate remained about 80 per cent. This device has been used to support patients in shock during coronary arteriography and left ventriculography in evaluation for myocardial revascularization. All 44 studied under these circumstances at the Massachusetts General Hospital had stenosis of the left anterior descending artery. The right coronary was obstructed in 77 per cent and triple vessel disease observed in 41 per cent. Akinesis or dyskinesis of the anterolateral wall of the left ventricle was demonstrated and when associated with no

perfusion, revascularization surgery was unsuccessful. Eight patients of 22 (36 per cent) with some visible arterial perfusion of akinetic regions survived aortocoronary bypass procedures.

Only one angiographically related death in this series of 44 high risk patients indicates the feasibility of assessment for surgery under these conditions. Those with destruction of more than 40 per cent of the left ventricle and poor regional arterial perfusion are unlikely to survive operation. Immediate survival of patients with less necrosis may be promising but long term benefits are not yet established.

Evaluation of the patient with acute myocardial infarction and shock requiring mechanical circulatory assistance and consideration for bypass grafting seem reasonable in this group with such a grave prognosis.

Left main coronary artery obstruction. Significant left main coronary stenosis places both anterior descending and circumflex branches in jeopardy and may compromise almost the total blood supply of the left ventricle. In most instances this is not an isolated lesion but is accompanied by double and triple vessel disease in the proximal left branch vessels and right coronary artery.

A higher risk of coronary arteriography has been reported with a 10 to 15 per cent mortality rate. Caution in study has been advised in patients with severe angina, strongly positive stress tests or calcification in the region of the left main trunk. Visualization of the right coronary first may demonstrate collateral filling of the left coronary system. Then a non-selective left aortic sinus injection of contrast medium may adequately delineate the left main obstruction without the dangers of compromise of the lumen by selective introduction of the catheter into the coronary ostium. This will be important to recognize lesions of the ostium which may not be appreciated with the catheter tip already within the coronary arterial lumen. Finally, cautious selective left coronary arteriograms can be accomplished for more detailed analysis.

There is a higher surgical risk in this group averaging about a 12 per cent mortality rate. A poor prognosis.

therapy and the presence of severe angina in most of the patients would seem to warrant consideration for early operation. This lesion is not common, occurring in less than 3 per cent of patients undergoing coronary arteriography and experience with surgery is not extensive. Further objective evaluation is necessary, particularly since left main arterial stenosis is almost never a solitary occlusive process.

Asymptomatic patients In general asymptomatic patients, despite documented coronary occlusive disease, are not operated upon except under certain special circumstances. Coronary arteriography in the assessment of valvular aortic stenosis may disclose morphologically significant obstructions in patients without angina. Coronary bypass is not performed routinely unless there is difficulty in weaning the patient off the pump oxygenator after prosthetic replacement of the valve or there is severe compromise of the coronaries with poor ventricular perfusion and anticipated low output failure in the early postoperative period. Similar situations may arise during ventricular aneurysmectomy and if feasible revascularization using one or more grafts may be attempted.

There is inadequate information as regards the benefits of surgery in the asymptomatic patient with uncomplicated prior myocardial infarction or demonstrable limited coronary artery disease without infarction. Prediction of impending myocardial infarction or sudden death in the asymptomatic patient is not yet possible. As high risk factors are identified in the natural course of the disease and further correlations are made with coronary arteriography and left ventricular function studies, perhaps a more liberal selectivity of candidates for surgery will follow.

Risk factors Hypertension, smoking, hypercholesterolemia and hyperlipidemia have been associated with acceleration of coronary atherosclerosis. The overall yearly mortality rate of 3 to 4 per cent in patients with angina is increased to 8 per cent in the subgroup with high blood pressure and advanced electrocardiographic abnormalities. Cardiomegaly and congestive heart failure adversely affect the prognosis.

With coronary arteriography we can attempt correlation of survival with dis-

tribution and extent of arterial stenosis. Multiple vessel disease, particularly involving the left anterior descending branch, worsens the prognosis. Approximately 90 per cent of patients with single vessel disease have a two year survival, only 65 to 75 per cent with two and three vessel involvement live that long. The importance of coronary collateral circulation and progression of obstructive lesions are variables with insufficient data for meaningful evaluation at this time.

As increase in vascular stenoses and reduction in left ventricular function influence the natural history of coronary arterial disease, these same factors increase the operative risks as well. The intraoperative and postoperative mortality rate and infarction rate are higher in those patients requiring multiple bypass grafts and in those with ventricular dysfunction.

In addition to the surgical mortality rate ranging from 3 to 10 per cent, we must consider the effectiveness of operation. The patency of saphenous vein aortocoronary grafts is in the range of 80 per cent after two years. A one year patency of 97 per cent has been claimed for internal mammary artery bypass, a procedure requiring longer operating time and with an obviously limited supply of vessels for grafting.

Conclusions

Adequate, statistically significant randomized data is not yet available to determine the effect of aortocoronary bypass grafting on the natural course of occlusive coronary atherosclerosis. Large prospective studies under way or in the planning stage may help define the influence of surgery upon future myocardial infarction and longevity.

There is no doubt of the relief of angina in about 85 per cent of patients undergoing operation. In the absence of left ventricular failure, operative mortality rate is low and if angina is incapacitating and unresponsive to medical management, surgery should be considered. Left main coronary lesions, unstable angina with acute unrelieved exacerbation and myocardial infarction with cardiogenic shock are high risk situations in which operation, if feasible, would be recommended.

Bypass surgery has been disappointing

in the treatment of chronic congestive heart failure. On the other side of the spectrum, operation has not been proved better than medical observation in the asymptomatic patient with old myocardial infarction. Acute uncomplicated myocardial necrosis will not be reversed by surgery, except possibly in the few special circumstances of coronary occlusion during coronary arteriography. In the light of our present knowledge there is little justification for aortocoronary bypass surgery in these three groups.

As our experience with coronary arteriography and surgery increases, perhaps we may liberalize the indications for operation. We must continue our objective evaluation of this new therapeutic modality and not finalize its role in the management of occlusive coronary atherosclerosis until its effects upon the natural history of the disease are established.

SUGGESTED READING

- Zukel W J, Cohen B M, Mattingly T W and Hrubec Z. Survey of following first diagnosis of coronary heart disease. *Am Heart J* 78:159 1969.
- Frank C W. The course of coronary heart disease: factors relating to prognosis. *Bull N Y Acad Med* 44:900 1968.
- Fulton M, Lutz W, Donald K W, Kirby B J, Duncan B, Morrison S L, Kerr I, Julian D G and Oliver M F. Natural history of unstable angina. *Lancet* i:860 1972.
- Oberman A, Jones W M, Riley C P, Reeves T J, Sheffield L T and Turner M E. Natural history of coronary artery disease. *Bull N Y Acad Med* 48:1109 1972.
- Irvinger G C, Ige F E and Ro S P. Prognostic significance of coronary arteriography. *Trans Assoc Am Physicians* 93:18 1970.
- Krass R R, Hutter A M Jr and DeSanctis R W. Acute coronary insufficiency: course and follow up. *Arch Intern Med* 129:808 1972.
- Favaloro R. Direct and indirect coronary surgery. *Circulation* 46:1197 1972.
- Manley J C and Johnson W D. Effects of surgery on angina (pre and post infarction) and myocardial function (failure). *Circulation* 46:1208 1972.
- Spencer I C, Green G E, Tice D A, Wallb E, Mills N I and Glassman I. Coronary artery bypass grafts for congestive heart failure: a report of experience with 40 patients. *J Thorac Cardiovasc Surg* 62:1529 1971.
- Irving P, Kimbiris D, Cecil B I and Linhart J W. Left main coronary artery disease. Clinical arteriographic and hemodynamic appraisal. *Am J Cardiol* 30:191 1972.
- Cohen M A, Cohn L I, Herman M A and Gorlin R. Diagnosis and prognosis of acute left coronary artery obstruction. *Circulation* 46 (Suppl 1):57 1972.
- Spencer I C. Bypass grafting for preinfarction angina. *Circulation* 45:1314 1972.
- Sinder C A, Buckley M J, Lemlich R C, Mundth J D and Lu ten W G. Mechanical circulatory assistance: Current status and experience with combining circulatory assistance emergency coronary angiography and acute myocardial revascularization. *Circulation* 46:1291 1972.

Annotations

Study of man, himself

No other animal or living thing has been studied as extensively as man. Man is intensely interested in himself, be he normal or diseased. However, in spite of extensive investigation of many sorts, there remain much to be learned about man by man. To know more about man, investigations must be conducted directly on man. Extrapolations from studies of other animals to man are not sufficient. The study of plant and other animals is extremely important and absolutely necessary even for a better understanding of man and his reactions. Nevertheless, investigation must be conducted on man himself. For example, diseases peculiar to man such as sickle cell anemia, certain inherited metabolic diseases, an inborn pectoral phoria, and other dermatologic states, obstetrical problems, allergic diseases such as asthma, psychiatric states such as schizophrenia, neuroathemia, urologic states such as enuresis, prostatism, and many others too numerous to even attempt to list must be studied on man.

Realizing that there is a need to study man, efforts must be made to avoid injury to man under study. Therefore, committees have been established to make certain man is not injured when subjected to research. This is good. However, it is unfortunate for many reasons that a committee for review of research plans for the study of man is necessary. A committee can have beneficial or detrimental influences on the study of man. And unfortunately too often some members of the committee have had absolutely no experience or very little experience with research on man.

Also, too often it is not fully realized that it is the investigator who really counts when considering the quality of the research and the subject's safety. A good, considerate, thoughtful, and humane physician will not subject a person to investigations which are hazardous. An investigator's attitude and approach to research on man can be judged best by his patient performance, including highly technical and scientific research on man as well as the use of new, yet untested drugs and procedure, diagnostic

and therapeutic (medical and surgical). Whether or not it is ethical or proper to charge a fee to those who are subjected to new and nonestablished diagnostic or therapeutic procedures is questionable. I think it is not proper. Nevertheless, at one time or another these various, often hazardous procedures and new drugs must be studied in man. Again, what is expected to happen and what actually happens are dependent entirely on the person responsible for the study.

Those who are primarily interested in man and who have devoted considerable effort in studying man know how delightful and satisfying it is to study man. The mere fact that the investigator can communicate directly with his subject during investigations offers tremendous advantages in obtaining data during the course of investigations. This is readily realized when studying pain. Man is the best animal to study in the conscious state. This type of study can seldom be done in other animals since the animals usually must be anesthetized merely to keep them quiet for recordings. Indeed, the anesthesia introduces an important and unknown variable. The psychic state of even conscious animals is difficult if not impossible to measure.

The human subject, when handled properly by the investigator, enjoys being studied and willingly participates, making the study his contribution to knowledge and for the improvement of the welfare of mankind. Man must be studied to obtain direct answers concerning man; he must be studied directly. But the researchee's welfare in every respect depends upon the researcher. It is a tremendous and invaluable privilege for the researcher to be permitted to study man. Let us not abuse that privilege under any circumstances. And remember there is no need for haste: studies on man must be safe.

G. E. Burch, M.D.
Department of Medicine
Tulane University School of Medicine
New Orleans, La.

Longevity of athletes

The confusing statistics on the longevity of athletes can probably be better understood by recognizing the different somatotypes of the athletes involved in the various sports.

Athletes come in all sizes and shapes filling every band of the somatotype spectrum from the ecto-

morphic marathoner (the average height and weight of 75 winners of the Boston Marathon is 5 ft 6 in and 130 pounds) to the endomorphic channel swimmer (who may attain 40 per cent body fat). Along this continuum you find the ectomesomorphic basketball players, move on to crewmen, and then

on to the endomorphous football players and weight lifters. Further into the endomorphic scale are baseball players and golfers.

Studies on the general population have already indicated that certain somatotypes especially the endomorphs have a shorter life span in our present culture. Spruin and co-workers³ as well as the Framingham results⁴ point toward this type's constitutional susceptibility to what Björck⁵ has called the disease of our century: coronary heart disease.

This may be genetically programmed or may simply be that the reactions to stress of the endomorphic—the socialization—and the mesomorphic—the action—are hazardous in present day society. The twin dangers of lipid affluence and an equilibriumism which allows no excuse for failure provide a perfect cultural medium for the endomorphic tendency to overindulge and the mesomorphic appetite for aggressive action.

The ectomorph who leans to withdrawal and asceticism tends to live longer. In times past he was more susceptible to tuberculosis and infection and his life span was therefore short. But present day killers he handles well.

Given this information one can understand why Prout⁶ found that oarsmen live more than six years longer than randomly selected classmates, why Karvonen's⁷ cross-country skiers outlived the national average by seven or more years.

One can also realize why Schnorr⁸ lumping all Danish champions together discovered that having attained the age of 50 years they merely lived out the normal Danish life span.

And one can see clearly why Largey⁹ scanning through Who Was Who in American Sports discovered trackmen lived longer by 14 years than football players—a finding confirmed by Polednack's¹⁰ extensive survey of 6,303 Harvard students.

Winning a letter might add years to your life reported Polednack—but only if it was in a minor sport. Major sport athletes died significantly earlier than non-athletes from coronary heart disease and perhaps earlier and oftener from tumors. All this information then falls into place. The differences detected by Prout⁶, Schnorr⁸, Largey⁹ and Polednack¹⁰ are simply the tendency of the population to have different life spans based on the reaction of their body builds with their environment.

The muscularly aggressive ex-football players are statistically susceptible to cardiovascular disease to just that degree shared by their muscular aggressive

non-football playing counterparts. Onlookers who share in the magnificent physical and psychological attributes of crewmen will share in their extended life span. Thin small boned people represented in sports by distance runners are longer lived than the average for regions already cited.

None of the studies incidentally claim any in formation on continuing athletic activity. It would be best to assume that there was little or none. Studies in England¹¹ a much more athletically inclined country disclosed that only 10 per cent of married men aged 23 to 30 continued with regular activity.¹²

We now know that ex-athletes seem to live as long as and no longer than non-athletes of the same body build.

What we need to know is whether athletic activity continuing through middle age will modify or maximize this genetically programmed longevity quotient. Will the perennial athlete outlive the ex-athlete of the same somatotype that?

George J. Sheehan MD
79 W. Front St.
Red Bank A J 08101

REFERENCES

1. Carter J E. The somatotypes of athletes: a review. *Hum Biol* 42:535 1970.
2. Spruin D M, Nathan D F and Gelles M. Weight, body type and the prevalence of coronary atherosclerotic heart disease in male. *Am J Med Sci* 215:63 1963.
3. Damon A et al. Predicting coronary heart disease from body measurements of Framingham males. *J Chronic Dis* 21:781 1969.
4. Björck G. Report of symposium on society stress and disease. Stockholm 1972.
5. Prout C. Life expectancy of college oarsmen. *JAMA* 220:1709 1972.
6. Karvonen M J. Report to international congress of gerontology 1969.
7. Schnorr P. Longevity and causes of death in male athletic champions. *Lancet* ii:1364 1971.
8. Largey D. Athletic activity and longevity. *Lancet* ii:286 1972.
9. Polednack A I. Longevity and cause of death among Harvard College athletes and their classmates. *Geriatrics* 27:53 1972.
10. Howell D H and Denis H I. The meaning of physical fitness: introduction. *Proc R Soc Med* 62:1155 1969.

Dopamine test for the diagnosis of coronary insufficiency

The limits of the exercise test for the diagnosis of coronary insufficiency are well known. We¹ have recently given evidence that infusion of isoproterenol is particularly useful for the diagnosis of coronary insufficiency in the absence of electrocardiographic (ECG) signs of the disease at rest. Similar results

have been obtained by Wexler, Kunitz, and Simonson.²

Better results have been obtained by our group with dopamine.³ In contrast to isopropyl norepinephrine, dopamine has very little effect on the heart rate; furthermore, at the dosage that we used it

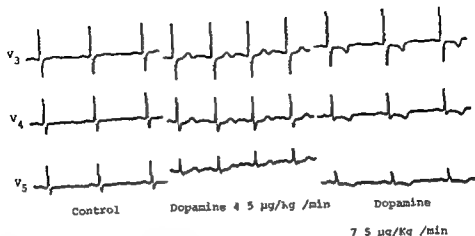


Fig 1 Effects of dopamine infusion on the ECG in a case of coronary artery disease. There is an evident initial transient improvement which is followed by ischemic changes of the ST segment.

does not increase the ventricular excitability and the peripheral vascular resistance.

As we know its primary cardiac effect is an increase in cardiac contractility and cardiac output; secondarily it increases the myocardial consumption of oxygen and the coronary blood flow.⁴

Our method is to use an intravenous administration of dopamine at an initial infusion rate of 4.5 µg per kilogram of body weight per minute. The infusion rate is then raised 1.5 µg per kilogram of body weight per minute every five minutes until a systolic blood pressure 50 per cent higher than the control value is achieved.

The test is stopped immediately in case subjective symptoms appear or if there are ECG signs of coronary insufficiency. The dopamine test has been compared with the cycloergometer test in supine position with increasing work load. Three groups of subjects have been studied. The first group consisted of 49 volunteers free of evidence of cardiac disease; in this group dopamine did not produce an ischemic depression of the S-T segment. At high dosages a junctional depression of the S-T segment and an increase in the voltage of T and U waves were observed almost constantly.

The second group consisted of 62 subjects with typical angina pectoris. The exercise test caused ischemic depression of the S-T segment in 61.2 per cent of the cases, while dopamine had the same effect in 72.6 per cent of the cases. The third group consisted of 45 patients with atypical precordial pain. The exercise test was positive in 20.0 per cent of the cases, while the dopamine test was positive in 28.9 per cent.

During the infusion of dopamine atrial ectopic beats were rarely observed in control subjects; in other subjects ventricular ectopic beats were observed in five cases; in one case a bigeminy appeared for a short time. The test is harmless with little subjective discomfort. Anginal pains were not more frequent with the dopamine test than with the exercise test.

An interesting observation is that the ischemic

ECG signs frequently appeared after the end of the dopamine infusion. This might be due to the fact that dopamine induces ischemic effects after a certain latency. An alternative hypothesis is its action on ionic exchanges which might hide the ischemic effects on the ECG. This is suggested by the results in other patients who at rest showed ischemic changes of the ECG. In these cases dopamine frequently caused surprising temporary changes in the shape of the ST segment and T wave toward an almost normal morphology. These changes usually appear before the accentuation of basal ischemic damage (Fig 1).

O Visoli MD
F N Effendy MD
G Malagnino MD†
Istituto di Clinica Medica I
Cardiovascular Section
University of Parma
43100 Parma Italy

†D. Malagnino died since the writing of this paper.

REFERENCES

- 1 Malagnino G, Astorri E, Chizzola A, Effendy F, N. Majorano C and Visoli O. Studio comparativo dello sforzo al cicloergometro e dell'infusione endovenosa di isopropilnoradrenalina nella diagnosi di insufficienza coronarica. *Ital Cardiol* 1:422 1971.
- 2 Wexler H, Kautz J and Simonson E. Electrocardiographic effects of isoprenaline in normal subjects and patients with coronary atherosclerosis. *Br Heart J* 33:759 1971.
- 3 Malagnino G, Effendy F, N. Astorri E, Manca C and Visoli O. Dopamine infusion test in the evaluation of coronary insufficiency. Comparison with exercise test. *G Ital Cardiol* 23:392 1972.
- 4 Goldberg L I. Cardiovascular and renal actions of dopamine: potential clinical applications. *Pharmacol Rev* 24:1 1972.

on to the endomorphoic football players and weight lifters. Further into the endomorphoic scale are baseball players and golfers.

Studies on the general population have already indicated that certain somatotypes especially the endomorphs have a shorter life span in our present culture. Spain and co-workers' findings² as well as the Framingham results³ point toward this type's constitutional susceptibility to what Björck⁴ has called the disease of our century: coronary heart disease.

This may be genetically programmed or may simply be that the reactions to stress of the endomorph—the socialization—and the mesomorph—the reaction—are hazardous in present day society. The twin dangers of lipid affluence and an egalitarianism which allows no excuse for failure provide a perfect cultural medium for the endomorphoic tendency to overindulge and the mesomorphoic appetite for aggressive action.

The ectomorph who leans to withdrawal and asceticism tends to live longer. In times past he was more susceptible to tuberculosis and infection and his life span was therefore short. But present day killers he handles well.

Given this information one can understand why Prout⁵ found that swimmers live more than six years longer than randomly selected classmates, why Karvonen's⁶ cross country skiers outlived the national average by seven or more years.

One can also realize why Schnorr⁷ lumping all Danish champions together discovered that having attained the age of 50 years they merely lived out the normal Danish life span.

And one can see clearly why Largey's⁸ scanning through Who Was Who in American Sports discovered trackmen lived longer by 14 years than football players, a finding confirmed by Polednick's⁹ extensive survey of 6,303 Harvard students.

Winning a letter might add years to your life reported Polednick—but only if it was in a minor sport.⁹ Major sport athletes died significantly earlier than non athletes from coronary heart disease and perhaps earlier and oftener from tumors. All this information then falls into place. The differences detected by Prout,⁵ Schnorr,⁷ Largey,⁸ and Polednick⁹ are simply the tendency of the population to have different life spans based on the reaction of their body builds with their environment.

The muscularly aggressive ex football player is statistically susceptible to cardiovascular disease to just that degree shared by their muscular aggressive

non football playing counterparts. Onlookers who share in the magnificent physical and psychological attributes of creammen will share in their extended life span. Thin, well boned people represented in sports by distance runners are longer lived than the average for reasons already cited.

None of the studies incidentally eluminate information on continuing athletic activity. It would be best to assume that there was little or none. Studies in England¹ of much more athletically inclined country disclosed that only 10 per cent of married men aged 23 to 30 continued with regular activity.¹¹

We now know that ex athletes seem to live as long as and no longer than non athletes of the same body build.

What we need to know is whether athletic activity continuing through middle age will modify or maximize this genetically programmed longevity quotient. Will the perennial athlete outlive the ex athlete of the same somatotype that is?

George I. Sheehan, MD
79 1/2 Front St.
Red Bank, N.J. 08041

REFERENCES

1. Carter J. L. The somatotypes of athletes: a review. *Hum Biol* 42:535 1970
2. Spain D. M., Nathan D. F. and Gelis M. Weight, body type and the prevalence of coronary atherosclerotic heart disease in male. *Am J Med Sci* 215:63 1963
3. Damon A. et al. Predicting coronary heart disease from body measurements of Framingham males. *J Chronic Dis* 21:781 1969
4. Björck G. Report of symposium on society stress and disease. Stockholm 1972
5. Prout C. Life expectancy of college oar men. *JAMA* 220:1709 1972
6. Karvonen M. J. Report to international congress of gerontology 1969
7. Schnorr I. Longevity and causes of death in male athletic champions. *Lancet* 11:1364 1971
8. Largey D. Athletic activity and longevity. *Lancet* 11:286 1972
9. Polednick A. P. Longevity and cause of death among Harvard College athletes and their classmates. *Geriatrics* 27:53 1972
10. Howell D. H. and Denis H. I. The measuring of physical fitness: introduction. *Proc R Soc Med* 62:1155 1969

Dopamine test for the diagnosis of coronary insufficiency

The limits of the exercise test for the diagnosis of coronary insufficiency are well known. We¹ have recently given evidence that infusion of isoproterenol is particularly useful for the diagnosis of coronary insufficiency in the absence of electrocardiographic (ECG) signs of the disease at rest. Similar results

have been obtained by Wexler, Hurty, and Simonson.²

Better results have been obtained by our group with dopamine.³ In contrast to isoproterenol, norepinephrine, dopamine has very little effect on the heart rate; furthermore, at the dosage that we used it

Letters to the Editor

Pacemaker catheter displacement

To the Editor

The recent report by Dr Preston on Electrocardiographic diagnosis of pacemaker catheter displacement in the JOURNAL (AM HEART J 85 445 1973) provides a simple way of such detection. However as can be seen from Figs 2 and 4 of the article the stimulus artifact is not the *only* thing that changes direction: the marked left axis deviation of the QRS in Fig 2 is no longer evident in Fig 4 (the precordial leads are also different). It would be of interest to know how many of Dr Preston's patients with catheter displacement showed a significant change in QRS and how many with stimulus artifact change showed a change in QRS.

Ludwig Klein MD
10 East 61th St
New York N Y 10021

electrocardiograms were shown. Also in Case 3 (Table I) the catheter displacement produced a clear change in the QRS vector and in Cases 5 and 7 the displacement was into the right atrium with subsequent loss of ventricular pacing.

I agree with Dr Klein's inference that catheter displacement can be diagnosed by changes in the QRS configuration as well as changes in the pacer artifact vector. When the catheter is displaced into the right ventricle however there is no pacer-induced QRS with which to make a comparison and in writing the article I chose not to discuss QRS changes so as to emphasize examination of the pacemaker artifact vector.

Thomas A Preston MD
Co Director Division of Cardiology
Harborview Medical Center
325 Ninth Ave
Seattle Wash 98104

Reply

To the Editor

Dr Klein correctly points out that the mean QRS vector also changed in the one case for which full

Complications of transfemoral coronary arteriography and their prevention using heparin

Coronary arteriography using the percutaneous femoral (Judkins) approach was introduced five years ago but only recently have reports appeared stressing the distressing complication rate. Chahine and colleagues¹ reported a 5 per cent incidence of myocardial infarction, stroke, or sudden death. Takaro and associates² reviewed 1,800 procedures from 20 Veterans hospitals and reported a 2.4 per cent fatality rate, attributing the majority of these deaths to thromboemboli.³ Green and co-workers³ reported 20 complications in 445 cases. These included five femoral thromboemboli, one of which was fatal.

In November 1970 those using the five catheterization laboratories in Seattle met informally to share their experience. There were 17 deaths associated with approximately 900 procedures over a three year period. Analysis of the 184 cases from the Providence Heart Center revealed 16 major complications with six deaths. Despite the liberal use of heparinized saline, the high incidence of thromboembolism was apparent. Furthermore, no learning curve was evident with the complications being evenly spread over three years. At that time I reported eight months' experience using heparin 50 mg immediately following the arterial puncture and an equal quantity of protamine sulfate at the end of the procedure. Thromboembolic complications stopped abruptly. This initial experience now extends to nearly three years and over 200 procedures still without a death or a major or minor thromboembolic complication. Furthermore, there have been no complications as a direct result of using heparin (currently we use 80 to 100 mg). Nachnani and co-workers⁴ described submicroscopic thrombi on polyurethane catheters despite the use

of heparin, but if this occurs clinically, it is apparently insignificant.

Experience at three Seattle hospitals (Providence, Swedish and Virginia Mason) now includes over 1,000 coronary arteriograms using total heparinization without fatality or thromboembolic complication.

The Seattle experience demonstrates the safety and efficacy of a single large dose of heparin before left ventriculography and selective coronary arteriography by way of the transfemoral approach and suggests that this method be employed until non thrombogenic catheters and guide wires become available.

Kenneth M Eyer MD
515 Minor Ave
Seattle Wash 98104

REFERENCES

- 1 Chahine R A, Herman M V and Gorlin R. Complications of coronary arteriography: comparison of the brachial to the femoral approach. *Ann Intern Med* 76:862 1972.
- 2 Takaro Timothy et al. Acute coronary occlusion following coronary arteriography: mechanisms and surgical relief. *Surgery* 72:1018 1972.
- 3 Green G S, McKinnon C M, Rosch J and Judkins M P. Complications of selective percutaneous transfemoral coronary arteriography and their prevention: review of 445 consecutive examinations. *Circulation* 45:552 1972.
- 4 Nachnani G H et al. Scanning electron microscopy of thrombogenesis on vascular catheter surfaces. *N Engl J Med* 286:139 1972.

Editorial

Central venous pressure: Physiological significance and clinical implications*

Arthur C Guyton M D
Carl E Jones Ph D
Jackson Miss

With the recent advent of continuous patient monitoring a factor frequently recorded is the central venous pressure. There is little doubt that knowing the level of central venous pressure and its changes can be of life saving value in some situations such as in the immediate post operative period following some types of cardiac surgery. On the other hand central venous pressure measurements have been extended to other clinical situations without as much reason and occasionally are used as a routine procedure in intensive care units or in monitoring programs. This has expanded considerably with the development of bedside catheterization techniques. The goal of the present editorial therefore is to bring into focus the physiological significance of central venous pressure its value as a clinical measurement and the relative advisability of making central venous pressure measurements in view of potential hazards particularly when made as a routine procedure.

When one speaks of central venous pressure he ideally means the pressure in the right atrium. However the pressure in the large systemic veins inside the chest of a supine person is almost invariably within

1 mm Hg of the pressure in the right atrium because the negative pressure of the thorax keeps these large veins in a distended state so that resistance to blood flow from them to the right atrium is extremely slight.^{1,2} Therefore central venous pressure can normally be measured with accuracy when the tip of the catheter is placed either into any of the great systemic veins of the thorax or into the right atrium itself.

Physiological significance of central venous pressure

Right atrial pressure (central venous pressure) has two major physiological effects. First it is this pressure that promotes filling of the heart during diastole and thereby plays a major role in determining the amount of blood pumped by the heart each minute—that is the level of cardiac output.^{4,7} For instance a 1 mm Hg increase in right atrial pressure can at times increase the cardiac output as much as 50 per cent and several mm Hg increase can cause as much as a threefold increase in cardiac output. Therefore it is immediately obvious that even slight changes in central venous pressure are significant in the control of over all circulatory adequacy.

From the Department of Physiology and Biophysics, University of Mississippi Medical Center, Jackson, Miss.
Received for publication Sept. 15, 1972.

Reprint requests to Arthur C. Guyton, M.D., Dept. of Physiology and Biophysics, The University of Mississippi Medical Center, 2500 N. State St., Jackson, Miss. 39216.

*The original investigations quoted in this article were supported by grants-in-aid from the United States Public Health Service.

Book reviews

✓ **CARDIOVASCULAR CLINICS** vol 4 No 3 Pediatric Cardiology Mary Allen Engle M D Guest editor Philadelphia 1972 F A Davis Company 366 pp Price \$12 00

This volume of Cardiovascular Clinics on Pediatric Cardiology is another excellent practical publication Mary Allen Engle has rendered an excellent service to cardiology She has gathered subjects of importance written by capable contributors The book is written not only for the pediatric cardiologist but for all physicians who treat heart disease in children The many subjects discussed are well selected and clearly presented This is another excellent addition to the series on Cardiovascular Clinics Dr Engle has done a fine job

✓ **VECTORCARDIOGRAPHY** By Alberto Benchimol M D Baltimore 1973 The Williams & Wilkins Company 223 pp Price \$18 50

Benchimol's book on vectorcardiography briefly reviews the principle of vectorcardiography lead

systems and its application in clinical medicine The selected systems discussed and the Frank system generally used are not clinically practical nor do any of them correct as is so often claimed the many variables that influence the resultant recording in all people or in the same person at all times The Frank system is difficult to apply in infants very sick patients and patients with chest deformities as well as difficult to repeat precise electrode placements at all times Until a simple practical method is introduced for general use such as the tetradral system vectorcardiography will never succeed as well as electrocardiography in clinical cardiology This book does present the VCG manifestations in selected common cardiac illnesses very well and is a useful book for beginners The field of course is much more extensive Therefore to become an expert in this field a study of the medical literature and a great deal of experience are necessary This book is a good one and is written for clinical application

Announcements

Pan Pacific Surgical Association Congress

The Pan Pacific Surgical Association announces its Thirteenth Congress to be held February 15 through 21 1975 in Honolulu Hawaii The Congress will convene at the Hilton Hawaiian Village Hotel and concurrent meetings will be held in anesthesiology colon and anorectal surgery general surgery neurosurgery obstetrics and gynecology ophthalmology orthopedic surgery otolaryngology plastic surgery thoracic cardiovascular surgery and urology

For further details regarding the Congress please write Dr Cesar M Defesus Pan Pacific Surgical Association 236 Alexander Young Building Honolulu Hawaii 96813

1973 Pediatric Cardiology Examination

The next examination in Pediatric Cardiology will be given in the fall of 1973 in the following manner

The written examination will be given on October 2 1973 in four cities Atlanta Chicago Philadelphia and San Francisco This part of the examination will require a full day A subsequent oral examination will be held on October 18 and 19 1973 in Chicago Illinois

Several rules regarding these examinations require emphasis (1) The fee paid for the examination is valid for the 1973 examination only (2) In order to be eligible to take the oral portion of the examination the candidate will have to achieve a *qualifying* score on the written examination (Please note that a *qualifying* grade is not necessarily a *passing* grade It would be possible to qualify with a low passing grade on the written pass the oral and still average out to a failing *total* score Conversely it would be possible to score high enough on the oral portion to pull a low qualifying written grade up to a total passing level) (3) A qualifying score on the 1973 written portion is valid only for the 1973 oral portion Any candidate who does not take the 1973 oral portion or who does not achieve a total passing grade in 1973 must take the entire examination on any subsequent attempt (4) Candidates who do not qualify on the written examination or who do not pass the total examination must pay a new examination fee for any subsequent re-examination

After the fall of 1973 examination subsequent examinations will be administered every other year—i.e. 1975 1977 1979 etc Since there will be a two year hiatus between the 1973 and 1975 examinations the deadline for receipt of applications for the 1973 examination was extended until July 1 1973

Still another modifier of the principle that a damaged heart is associated with increased central venous pressure is the effect of neural stimulation on the heart. Even without neural stimulation the heart has the capacity to pump two to three times normal amounts of cardiac output without significant elevation of the central venous pressure.¹¹ Therefore slight damage to the heart rarely causes a noticeable change in central venous pressure. And even when cardiac damage becomes moderately extensive the central venous pressure is still likely to remain within normal limits because of the compensating effects of neural stimulation. For instance the fast heart rate and sometimes increased thrust of the heart in moderate heart disease are both manifestations of increased neural drive to the heart initiated by reflexes (1) from the heart itself,¹¹ (2) from baroreceptors in the great arteries,^{22,23} or occasionally (3) from chemosensitive areas in the brain.²⁴ Thus the increased neural drive to the heart can often compensate for diminished intrinsic cardiac performance and the result is often an insignificant change or no change in central venous pressure despite moderately severe cardiac debility.

Effect of tendency for venous return on central venous pressure. One can readily understand that when increased quantities of blood are attempting to return to the heart from the peripheral circulation this also will increase the central venous pressure. However if the heart is completely normal even doubling the venous return will cause only a 1 to 2 mm Hg increase in central venous pressure because the heart will simply pump the increased blood flowing into the right atrium.²⁵ Therefore in the face of a normal heart the tendency for venous return must increase vastly to raise central venous pressure significantly.

On the other hand if the pumping effectiveness of the heart has been severely diminished even the slightest increased tendency for venous return may engorge the central veins and the right atrium with so much blood that the central venous pressure will rise markedly. This is the effect that occurs in chronic congestive heart failure in which the heart itself is weak in addition to the greatly increased tendency

for venous return thus markedly elevating central venous pressure.

Causes of increased tendency for venous return. Two principal causes of increased tendency for venous return are (1) decreased resistance to blood flow from the arteries to the veins^{26,27} and (2) increased ratio of blood volume to vascular blood holding capacity.²⁸

An ultimate form of decreased peripheral resistance occurs in patients with large A-V fistulae in which case the tendency for venous return can often be increased even chronically to as much as two to three times normal.^{29,30} Yet here again if the heart has sufficient pumping effectiveness the excess blood returning to the heart will be pumped immediately on into the arteries so that the increase in central venous pressure may be as little as 1 mm Hg.

An increase in the ratio of blood volume to vascular blood holding capacity means simply that more blood is present in the vascular tree than the vascular tree is designed to hold. For instance let us assume that the blood volume is increased 50 per cent but that the vascular tree is still of normal basic size. To accommodate the increased volume all or most of the blood vessels of the body become considerably distended. As a result the pressures in all these vessels increase and these increased pressures increase the tendency for blood to flow from the peripheral vessels toward the low pressure area of the central veins.³¹ The greater the pressure differences between the peripheral vessels and the central veins the greater also will be the rate of venous return. Yet if the heart has normal pumping effectiveness it is very difficult for the venous return to rise high enough to cause more than minor increases in central venous pressure. Yet on the other hand the combination of a weakened heart and greatly increased blood volume can cause inordinately elevated central venous pressure.

Effect of external cardiac pressure on central venous pressure. Since the manometers used for measuring central venous pressure are themselves exposed to atmospheric pressure any change in pressure surrounding the heart and central veins will be reflected by a similar change in the pressure reading from the manometer. A classic example of this effect occurs in cardiac

The second reason for extreme significance of central venous pressure is that it acts as a back pressure on the systemic circulation to oppose return of blood from the peripheral blood vessels into the heart¹¹. Here again, relatively slight changes in central venous pressure can under some conditions have marked effects on the circulation. For instance, *in an animal whose cardiovascular reflexes have been blocked* an increase in central venous pressure from its normal value of approximately 0 mm Hg to a positive value of only 7 mm Hg will usually completely stop venous return.

Fortunately the extreme effects on both the heart and on venous return caused by changes in central venous pressure are greatly moderated by the circulatory neural reflexes¹¹. Therefore, under normal conditions the central venous pressure can change upward or downward several mm Hg without significantly altering blood flow in the circulatory periphery. And by the same token this means that measurements of the central venous pressure in the normal person can vary as much as 3 to 4 mm Hg without having real significance. Yet there are limits to function of the neural reflexes, so that central venous pressure changes beyond a few mm Hg do have serious effects on function of the circulation.

Normal and abnormal factors that alter central venous pressure

Two very obvious factors that affect central venous pressure are (1) pumping effectiveness of the heart and (2) tendency for venous return. However another factor that can alter central venous pressure tremendously and that is often forgotten is (3) the pressure surrounding the heart. These three different factors affect central venous pressures in different ways and they all have different physiological significances as follows.

Role of the heart in determining central venous pressure. One can readily understand that decreased effectiveness of the heart as a pump will tend to dam blood behind the heart and, therefore will tend to elevate central venous pressure¹²⁻¹⁵. However this simple viewpoint needs to be modified in several different ways.

First cardiac damage is not always associated with elevated central venous pressure. For instance acute damage to the left ventricle alone without significant damage to the right ventricle occasionally causes either no change in right atrial pressure or even a decrease¹²⁻¹⁴. The reason for this is that the associated pulmonary congestion causes large amounts of blood to translocate from the systemic circulation into the lungs. When this happens, too little blood is available in the systemic circulation for adequate return of blood through the systemic veins into the right atrium. Yet, at the same time the right heart is still pumping blood with almost usual effectiveness. Therefore in the face of an almost normal right heart but too little return of blood to the heart, one can readily understand why acute left ventricular failure is frequently associated with very little change in central venous pressure, sometimes with a slight rise sometimes with no change at all and occasionally with an actual decrease.

A second modification of the principle that a damaged heart causes increased central venous pressure is that chronic cardiac damage almost always increases the central venous pressure far more than does acute damage^{12-14,16}. For instance acute generalized damage of the heart that reduces the cardiac output almost to lethal levels is often associated with no more than a few mm Hg rise in central venous pressure. Yet as the condition proceeds into a chronic state the central venous pressure may rise as much as 15 to 20 mm Hg. This chronic effect is generally caused by renal fluid retention secondary to the cardiac debility with subsequent increase in blood volume. Initiating causes of the fluid retention are (1) slightly reduced arterial pressure which causes reduction in renal blood flow and glomerular filtration^{17,18} (2) an intense increase in sympathetic vasoconstriction of the renal arterioles which further reduces glomerular filtration and (3) an increased secretion of aldosterone which in turn promotes salt and water retention by the kidneys^{19,20}. Therefore in chronic cardiac insufficiency the increased central venous pressure is a product not only of reduced pumping effectiveness of the heart but also of increased tendency for venous return caused by excess body fluid and blood.

Still another modifier of the principle that a damaged heart is associated with increased central venous pressure is the effect of neural stimulation on the heart. Even without neural stimulation the heart has the capacity to pump two to three times normal amounts of cardiac output without significant elevation of the central venous pressure.¹¹ Therefore slight damage to the heart rarely causes a noticeable change in central venous pressure. And even when cardiac damage becomes moderately extensive the central venous pressure is still likely to remain within normal limits because of the compensating effects of neural stimulation. For instance the fast heart rate and sometimes increased thrust of the heart in moderate heart disease are both manifestations of increased neural drive to the heart initiated by reflexes (1) from the heart itself,¹² (2) from baroreceptors in the great arteries,¹³ or occasionally (3) from chemosensitive areas in the brain.¹⁴ Thus the increased neural drive to the heart can often compensate for diminished intrinsic cardiac performance and the result is often an insignificant change or no change in central venous pressure despite moderately severe cardiac debility.

Effect of tendency for venous return on central venous pressure. One can readily understand that when increased quantities of blood are attempting to return to the heart from the peripheral circulation this also will increase the central venous pressure. However if the heart is completely normal even doubling the venous return will cause only a 1 to 2 mm Hg increase in central venous pressure because the heart will simply pump the increased blood flowing into the right atrium.¹⁵ Therefore in the face of a normal heart the tendency for venous return must increase vastly to raise central venous pressure significantly.

On the other hand if the pumping effectiveness of the heart has been severely diminished even the slightest increased tendency for venous return may engorge the central veins and the right atrium with so much blood that the central venous pressure will rise markedly. This is the effect that occurs in chronic congestive heart failure in which the heart itself is weak in addition to the greatly increased tendency

for venous return thus markedly elevating central venous pressure.

Causes of increased tendency for venous return. Two principal causes of increased tendency for venous return are (1) decreased resistance to blood flow from the arteries to the veins,^{16,17} and (2) increased ratio of blood volume to vascular blood holding capacity.¹⁸

An ultimate form of decreased peripheral resistance occurs in patients with large A-V fistulae in which case the tendency for venous return can often be increased even chronically to as much as two to three times normal.^{19,20} Yet here again if the heart has sufficient pumping effectiveness the excess blood returning to the heart will be pumped immediately on into the arteries so that the increase in central venous pressure may be as little as 1 mm Hg.

An increase in the ratio of blood volume to vascular blood holding capacity means simply that more blood is present in the vascular tree than the vascular tree is designed to hold. For instance let us assume that the blood volume is increased 50 per cent but that the vascular tree is still of normal basic size. To accommodate the increased volume all or most of the blood vessels of the body become considerably distended. As a result the pressures in all these vessels increase and these increased pressures increase the tendency for blood to flow from the peripheral vessels toward the low pressure area of the central veins.²¹ The greater the pressure differences between the peripheral vessels and the central veins the greater also will be the rate of venous return. Yet if the heart has normal pumping effectiveness it is very difficult for the venous return to rise high enough to cause more than minor increases in central venous pressure. Yet on the other hand the combination of a weakened heart and greatly increased blood volume can cause inordinately elevated central venous pressure.

Effect of external cardiac pressure on central venous pressure. Since the manometers used for measuring central venous pressure are themselves exposed to atmospheric pressure any change in pressure surrounding the heart and central veins will be reflected by a similar change in the pressure reading from the manometer. A classic example of this effect occurs in cardiac

tamponade in which the pressure within the pericardial cavity is greatly increased. This increased pressure makes it impossible for the right atrium to fill with blood until the central venous pressure outside the pericardium rises to a level slightly above the pericardial pressure.^{11, 12} Yet, when the right atrium does fill adequately with blood, the blood can be pumped as usual by the heart. Therefore, even if the heart itself is normal, the central venous pressure will be increased an amount almost equal to the cardiac tamponade pressure. If in addition to the cardiac tamponade the pumping effectiveness of the heart is also reduced, this can cause still more rise in central venous pressure.

Though cardiac tamponade is a dramatic example of the effect of external cardiac pressure on central venous pressure, a much more common example is simply pressure changes related to respiration. With each breathing cycle the pressure surrounding the heart and great veins of the chest increases and decreases. Therefore the measured central venous pressure also rises and falls.^{13, 14} In addition, respiratory abnormalities can often cause chronic elevation or depression of venous pressure for instance:

- 1 If a patient develops bilateral pneumothorax or if the thorax is opened the pericardial pressure can rise from its normal value of -4 mm Hg up to a value approaching atmospheric pressure, thus elevating the central venous pressure by approximately the same amount.¹⁵

- 2 Positive pressure breathing by a patient increases the external cardiac pressure sometimes to many mm Hg positive pressure and, therefore, can also elevate the central venous pressure.¹⁶

- 3 Increased resistance to airflow in the lungs, such as in asthma or in some instances of emphysema, can often cause greatly increased positive intra-alveolar pressures during expiration, thereby seriously increasing the average external cardiac pressure. This pressure rise is also reflected in the central venous pressure.¹⁷

- 4 When patients overexpand their lungs, the increased stretch of the lungs increases the recoil pressure of the lungs, which in turn decreases the pressure surrounding the heart and results in a corresponding decrease in central venous pressure. There-

fore, simply changing the average volume of air in the lungs will significantly alter the central venous pressure.

In summary, external cardiac factors related either to the pericardium, as in cardiac tamponade, or to positive or negative pressures caused by normal or abnormal respiration can frequently alter the central venous pressure as much as 4 to 5 mm Hg and occasionally as much as 70 to 30 mm Hg. Since this variable is not always directly related to the adequacy of the circulation itself, it can often obscure the cardiac or peripheral circulatory factors that tend to alter the central venous pressure.

Hydrostatic pressure reference level for measuring central venous pressure

Another problem in assessing the significance of central venous pressure measurements is the hydrostatic pressure level to which the central venous pressure is referred. It has already been pointed out that even a millimeter or so pressure change can sometimes cause as much as 50 to 100 per cent change in cardiac output. If one wishes to assess the adequacy of input pressure to the heart, it is important that the pressure level within the right atrium be estimated accurately within 1 mm Hg. Therefore, what is the proper hydrostatic pressure level to which the central venous pressure should be referred? One of the most widely used reference levels is that of Winsor and Burch¹⁸ based on measurements in a large number of adults. They found the pressure to vary least from one normal supine person to another when the hydrostatic reference level was considered to be approximately the level of the posterior surface of the right atrium. In other hemodynamic studies in dogs, Guyton and Greganti¹⁹ showed that the right atrial pressure remains almost exactly constant in an animal rotated into any position in space provided that the hydrostatic reference point is the geometric center of the tricuspid valve. This point was found to be 61 per cent of the thickness of the chest in front of the animal's back and 77 per cent of the distance from the suprasternal notch to the tip of the xiphoid process.

Yet, regardless of the precise hydrostatic

reference point used by each investigator or clinician it is very clear that the absolute value of the central venous pressure measurement is no more accurate than the precision of the hydrostatic reference level to which the pressure is referred. On the other hand if one is looking simply for changes in central venous pressure in a single patient the hydrostatic reference level becomes of little importance.

Clinical value of central venous pressure measurements

The discussion thus far has presented two contrasting points of view: first the extreme importance of central venous pressure in determining both cardiac output and venous return; but second the many different factors that can alter the central venous pressure and thereby often make difficult the interpretation of central venous pressure measurements. The extreme importance of central venous pressure explains the desire of many clinicians to measure it or even to monitor it continuously. Yet on the other hand the difficulty of deciding which of the many determinants of central venous pressure might be causing an abnormal pressure mitigates against the usefulness of the measurements that are made. When one is almost certain which of the multiple factors is likely to alter the central venous pressure, a continuous record of the pressure is often very valuable for alerting one to a changing condition. For instance following acute cardiac surgery it is usually the heart that is the determining factor of changes in central venous pressure or it might be a combination of the heart and the blood volume if significant quantities of parenteral fluid are being administered. But at least the choices are few.

Another instance in which central venous pressure measurements seem to have been used to advantage is in the treatment of circulatory shock, particularly when the treatment involves administration of large volumes of fluid. For instance studies by Brand and associates²¹ and others have indicated the importance of keeping a slightly elevated central venous pressure during the course of fluid administration in severe degrees of circulatory shock. The slightly elevated central venous pressure can be particularly helpful in making the heart

pump a normal cardiac output even when the cardiac strength has become decreased as occurs in the late stages of circulatory shock.^{22, 23}

But now we come to the less acute and less life threatening situations in which one might wish to know the central venous pressure level but in which this is not a necessary component either of diagnosis or of therapy. For instance does one need to know the central venous pressure to diagnose congestive heart failure? In general he can estimate this within 1 to 2 mm Hg by measuring the peripheral venous pressure because the veins in severe congestive failure are so well distended that there is little pressure drop from the peripheral veins to the heart. Indeed a good clinician can estimate the central venous pressure within 2 to 3 mm Hg by simply observing the degree of distention of the neck veins in both the supine and sitting positions. And such clinical signs as skin color, skin temperature, pulse rate, cardiac thrust, the dyspnea index, presence or absence of pulmonary rales and so forth are usually much more directly to the point of the circulatory deficiency than is the central venous pressure anyway.

Thus it is with ambivalence that we have discussed central venous pressure in view of the clear and unmistakable importance of this pressure level to the function of the circulatory system, but also in view of the difficulty of interpretation and the possible hazards of the procedure itself. These hazards are generally well known but they are sufficiently important that they should be left in the mind of the reader: (1) possible clotting aftermaths; (2) possible internal damage to the blood vessels themselves or to the heart; (3) possible introduction of infection into the circulation; and (4) inadvertent initiation of cardiac arrhythmias—sometimes even cardiac fibrillation particularly when a catheter inadvertently passes into the right ventricle or when an electrical current is carried to the inner wall of the heart from the recording manometer through the catheter, a not too uncommon cause of death.

REFERENCES

1. Guyton A. C. and Adkins L. H. Quantitative aspects of the collapse factor in relation to ve-

- nous return *Am J Physiol* 177 523 1954
- 2 Guyton A C Lindsey A W Abernathy J B and Richardson T Q Venous return at various right atrial pressures and the normal venous return curve *Am J Physiol* 189 609 1957
- 3 Guyton A C Langston J B and Carrier O Decreased venous return caused by right atrial pulsation *Circ Res* 10 188 1962
- 4 Patterson S W and Starling E H On the mechanical factors which determine the output of the ventricles *J Physiol (Lond)* 48 357 1914
- 5 Sarnoff S J Myocardial contractility as described by ventricular function curves observations on Starling's Law of the Heart *Physiol Rev* 35 107 1955
- 6 Sagawa K Analysis of the ventricular pumping capacity as a function of input and output pressure loads in Reeve E B and Guyton A C editors *Physical bases of circulatory transport Regulation and exchange Philadelphia 1967 W B Saunders Company* p 141
- 7 Herndon C W and Sagawa K Combined effects of aortic and right atrial pressures on aortic flow *Am J Physiol* 217 65 1969
- 8 Guyton A C Venous return in Hamilton W F and Dow P editors *Handbook of physiology Circulation Washington D C 1963 American Physiological Society* p 1099
- 9 Guyton A C Lindsey A W Abernathy J B and Langston J B Mechanism of the increased venous return and cardiac output caused by epinephrine *Am J Physiol* 192 126 1958
- 10 Feroso J D Richardson T Q and Guyton A C Mechanism of decrease in cardiac output caused by opening the chest *Am J Physiol* 207 1112 1964
- 11 Richardson T Q Feroso J D and Pugh G O Effect of acutely elevated intracranial pressure on cardiac output and other circulatory factors *J Surg Res* 5 318 1965
- 12 Stone H L Bishop V S and Guyton A C Progressive changes in cardiovascular function after unilateral heart irradiation *Am J Physiol* 206:289 1964
- 13 Kenner H M and Wood E H Intrapericardial intrapleural and intracardiac pressures during acute heart failure in dogs studied with out thoracotomy *Circ Res* 19 1071 1966
- 14 Guyton A C Regulation of cardiac output *N Engl J Med* 277 805 1967
- 15 Dodge H T and Baxley W A Hemodynamic aspects of heart failure *Am J Cardiol* 22 74 1968
- 16 Baumber J S Davis J O Schneider E G and Johnson J A Plasma renin activity and the effects of deoxycorticosterone acetate in dogs with chronic left ventricular overload *Circ Res* 27:705 1970
- 17 Baer P G Navar L G and Guyton A C Renal autoregulation filtration rate and electrolyte excretion during vasodilatation *Am J Physiol* 219 619 1970
- 18 Navar L G Uther J B and Baer P G Pressure diuresis in dogs with diabetes *insipidus Nephron* 8 97 1971
- 19 Davis J O Kliman B Yarkopoulos A and Peterson R E Increased aldosterone secretion following acute constriction of the inferior vena cava *J Clin Invest* 31 193 1953
- 20 Davis J O Physiology of congestive heart failure in Hamilton W F and Dow P editors *Handbook of physiology sec. 7 vol. 2 Circulation Washington D C 1965 American Physiological Society* p 2093
- 21 Douthett U and Kramer K Über die druckregulierung kreislaufregulieren-der reflexe am linken herz *Pflügers Arch* 269 114 1959
- 22 Iriuchijima J Soulsby M E and Wilson M F Participation of cardiac sympathetic nerves in carotid occlusion pressor reflex *Am J Physiol* 215 1111 1968
- 23 Allison J L Sagawa K and Kumada M An open loop analysis of the aortic arch barostatic reflex *Am J Physiol* 217 1576 1969
- 24 Downing S E Mitchell J H and Wallace A C Cardiovascular responses to ischemia hypoxia and hypercapnia of the central nervous system *Am J Physiol* 204 681 1963
- 25 Guyton A C and Sagawa K Compensations of cardiac output and other circulatory functions in reflex dogs with large A V fistulae *Am J Physiol* 200 1157 1966
- 26 Guyton A C Determination of cardiac output by equating venous return curves with cardiac response curves *Physiol Rev* 35 123 1955
- 27 Guyton A C Abernathy J B Langston J B Kaufmann B N and Fairchild H M Relative importance of venous and arterial resistance in controlling venous return and cardiac output *Am J Physiol* 196 1008 1959
- 28 Guyton A C Lindsey A W and Kaufmann B Effect of mean circulatory filling pressure and other peripheral circulatory factors on cardiac output *Am J Physiol* 180 463 1955
- 29 Crawford E S Turell D J and Alexander J K Aorto-inferior vena caval fistula of neoplastic origin Hemodynamic and coronary blood flow studies *Circulation* 27 414 1963
- 30 Samet P Bernstein W H Jacobs W and Fomon J Indicator dilution curves in systemic arteriovenous fistulas *Am J Cardiol* 13 176 1964
- 31 Isaacs J P Berglund E and Sarnoff S J Ventricular function III The pathologic physiology of acute cardiac tamponade studied by means of ventricular function curves *Am HEART J* 48 66 1954
- 32 Fowler N O and Holmes J C Hemodynamic effects of isoproterenol and norepinephrine in acute cardiac tamponade *J Clin Invest* 48 502 1969
- 33 Abel F L and Waldhausen J A Respiratory and cardiac effects on venous return *Am HEART J* 73 286 1969
- 34 Morgan B C Guntheroth W G and Dillard O H Relationship of pericardial to pleural pressure during quiet respiration and cardiac tamponade *Circ Res* 16 493 1965

- 35 Morgan B C Martin W E Hornbein T F Crawford E W and Guntheroth W G Hemodynamic effects of intermittent positive pressure respiration *Anesthesiology* 27 584 1966
- 36 Winsor T and Burch G E Use of the phlebomanometer. Normal venous pressure values and a study of certain clinical aspects of venous hypertension in man *AM HEART J* 31 387 1946
- 37 Guyton A C and Griganti L P A physiologic reference point for measuring circulatory pressures in the dog—particularly venous pressure *Am J Physiol* 185 137 1956
- 38 Brand E O Eadie E H Feler R J Winkler C W and Goldsmith R H Metabolic vs cardiovascular dynamic support in therapy of posthemorrhagic shock *Am J Physiol* 220:1437 1971
- 39 Crowell J W and Guyton A C Further evidence favoring a cardiac mechanism in irreversible hemorrhagic shock *Am J Physiol* 203:248 1967
- 40 Bethea H L Jones C E and Crowell J W Effect of pharmacologic coronary flow augmentation on cardiac function in hypotension *Am J Physiol* 222 95 1972

- nous return *Am J Physiol* 177:523 1954
- 2 Guyton A C Lindsey A W Abernathy J B and Richardson T Q Venous return at various right atrial pressures and the normal venous return curve *Am J Physiol* 189:609 1957
- 3 Guyton A C Langston J B and Carrier O Decreased venous return caused by right atrial pulsation *Circ Res* 10:188 1962
- 4 Pitterson S W and Starling E H On the mechanical factors which determine the output of the ventricles *J Physiol (Lond)* 18:357, 1914
- 5 Sarnoff S J Myocardial contractility as described by ventricular function curves observations on Starlings Law of the Heart *Physiol Rev* 35:107 1955
- 6 Sagawa K Analysis of the ventricular pumping capacity as a function of input and output pressure loads in Reeve E B and Guyton A C editors *Physical bases of circulatory transport Regulation and exchange Philadelphia 1967 W B Saunders Company* p 141
- 7 Herndon C W and Sagawa K Combined effects of aortic and right atrial pressures on aortic flow *Am J Physiol* 217:65 1969
- 8 Guyton A C Venous return in Hamilton W F and Dow P editors *Handbook of physiology Circulation* Washington D C 1963 American Physiological Society p 1099
- 9 Guyton A C Lindsey A W Abernathy J B and Langston J B Mechanism of the increased venous return and cardiac output caused by epinephrine *Am J Physiol* 192:126 1958
- 10 Feroso J D Richardson T Q and Guyton A C Mechanism of decrease in cardiac output caused by opening the chest *Am J Physiol* 207:1112 1964
- 11 Richardson T Q Feroso J D and Pugh G O Effect of acutely elevated intracranial pressure on cardiac output and other circulatory factors *J Surg Res* 5:318 1965
- 12 Stone H L Bishop V S and Guyton A C Progressive changes in cardiovascular function after unilateral heart irradiation *Am J Physiol* 206:289 1964
- 13 Kenner H M and Wood E H Intrapericardial intrapleural and intracardiac pressures during acute heart failure in dogs studied with out thoracotomy *Circ Res* 19:1071 1966
- 14 Guyton A C Regulation of cardiac output *N Engl J Med* 277:805 1967
- 15 Dodge H T and Baxley W A Hemodynamic aspects of heart failure *Am J Cardiol* 22:24 1968
- 16 Baumber J S Davis J O Schneider E G and Johnson J A Plasma renin activity and the effects of deoxycorticosterone acetate in dogs with chronic left ventricular overload *Circ Res* 27:705 1970
- 17 Baer P G Navar L G and Guyton A C Renal autoregulation filtration rate and electrolyte excretion during vasodilatation *Am J Physiol* 219:619 1970
- 18 Navar L G Uther J B and Baer P G Pressure diuresis in dogs with diabetes insipidus *Nephron* 8:97 1971
- 19 Davis J O Klimm B Yankopoulos N A and Peterson R E Increased aldosterone secretion following acute constriction of the inferior vena cava *J Clin Invest* 3:158 1958
- 20 Davis J O Physiology of congestive heart failure in Hamilton W F and Dow P editors *Handbook of physiology* sec 2 vol 3 Circulation Washington D C 1965 American Physiological Society p 2095
- 21 Douthett U and Kramer K Über die druckregulierung kreislaufregulieren-der reflexe aus dem linken herz *Pflügers Arch* 269:114 1959
- 22 Iriuchijima J Soulsby M E and Wilson M F Participation of cardiac sympathetic in carotid occlusion pressor reflex *Am J Physiol* 215:1111 1968
- 23 Allison J L Sagawa K and Kumada M An open loop analysis of the aortic arch barostatic reflex *Am J Physiol* 217:1576 1969
- 24 Downing S E Mitchell J H and Wallace A C Cardiovascular responses to ischemia hypoxia and hypercapnia of the central nervous system *Am J Physiol* 204:881 1963
- 25 Guyton A C and Sagawa K Compensations of cardiac output and other circulatory functions in areflex dogs with large A V fistulas *Am J Physiol* 200:1157 1966
- 26 Guyton A C Determination of cardiac output by equating venous return curves with cardiac response curves *Physiol Rev* 35:173 1955
- 27 Guyton A C Abernathy J B Langston J B Kaufmann B N and Fairchild H M Relative importance of venous and arterial resistance in controlling venous return and cardiac output *Am J Physiol* 196:1008 1959
- 28 Guyton A C Lindsey A W and Kaufmann B Effect of mean circulatory filling pressure and other peripheral circulatory factors on cardiac output *Am J Physiol* 180:463 1955
- 29 Crawford E S Turell D J and Alexander J K Aorto-inferior vena caval fistula of neoplastic origin Hemodynamic and coronary blood flow studies *Circulation* 27:414 1963
- 30 Samet P Bernstein W H Jacobs W and Fomon J Indicator dilution curves in systemic arteriovenous fistulas *Am J Cardiol* 13:176 1964
- 31 Isaacs J P Berglund E and Sarnoff S J Ventricular function III The pathologic physiology of acute cardiac tamponade studied by means of ventricular function curves *Am Heart J* 48:66 1954
- 32 Fowler N O and Holmes J C Hemodynamic effects of isoproterenol and norepinephrine in acute cardiac tamponade *J Clin Invest* 48:502 1969
- 33 Abel F L and Waldhausen J A Respiratory and cardiac effects on venous return *Am Heart J* 71:266 1969
- 34 Morgan B C Guntheroth W G and Dillard O H Relationship of pericardial to pleural pressure during quiet respiration and cardiac tamponade *Circ Res* 16:493 1965



Fig 1 A and B The opacified left ventricle in end diastole (A left) and in end systole (B right) before operation

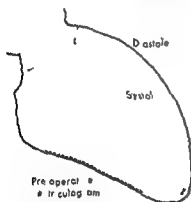


Fig 1C Superimposed tracings of the outline in both phase illustrate akinesis of the inferior wall and hypokinesia of the anterior and apical segments of the left ventricle

and true aneurysm were not included in this series. The walls of the left ventricle identified as anterior, apical and inferior were studied separately although most patients had more than one asynergic wall involvement or instances of asynergy. Only asynergic walls associated with patent grafts to corresponding coronary arteries were included in the study. Forty-eight such instances of ventricular wall asynergy were found in 37 patients and are described in Table I.

A standard twelve-lead ECG was recorded on a direct writing Sanborn machine. The technique of recording VCGs was that devised by Frank, using the fifth intercostal space with the patient in the sitting position. The recordings were done

Table I Type and site of instances of wall asynergy. The site refers to left ventricular areas visualized on ventriculograms obtained in the right oblique position.

| Type | Site of asynergy | | |
|-------------|------------------|------|----------|
| | Anterior | Apex | Inferior |
| Akinesis | 2 | 4 | 9 |
| Hypokinesis | 12 | 19 | 2 |

on a 1507A Vector Programmer (Hewlett Packard). The ECG and VCG were recorded on the same day. Criteria used for the diagnosis of transmural infarction by ECG and VCG were those which have been generally accepted.^{11,20}

The distribution of asynergy and myocardial infarction (MI) as diagnosed by ECG/VCG, excluding left ventricular hypertrophy (LVH) and left bundle branch block (LBBB) are shown in Table II. Infarctions were detected in the following areas: anterior, anterolateral, high lateral, anteroapical, inferior and true posterior and thus were studied separately. There were 23 instances of infarction diagnosed by ECG criteria and 24 instances detected by VCG criteria. Nine instances of infarction were excluded because they were not related to areas supplied by coronary arteries having patent grafts. Infarctions localized to the anterior, anterolateral, high lateral and anteroapical areas were related to

Improvement of left ventricular asynergy following aortocoronary bypass surgery related to preoperative electrocardiogram and vectorcardiogram

Kurian Jacob, MD

Michel Chabot, MD

Jacques Saltiel, MD

Lucien Campeau, MD

Montreal, Canada

In the last three years an increasing number of patients have had aorta to coronary saphenous bypass grafts.¹⁻⁶ Studies from this Institution^{7,8} have shown that localized disorder of left ventricular (LV) wall contraction or asynergy may be partially or totally corrected following this type of operation. The present study was undertaken to determine whether the electrocardium and vectorcardiogram (ECG/VCG) might predict the outcome of these anomalies of contraction postoperatively.

Materials and methods

Anomalies of ventricular contraction localized to the anterior apex and inferior areas of the left ventricle were studied separately. These anomalies of contraction were detected among 50 patients who were from the first group operated upon in this Institution. We have excluded subjects who had internal mammary artery implantation or muscle resection in addition to the bypass graft procedures. All these patients had complete preoperative studies, namely ECG, VCG, coronary arteriography and left ventriculography. Of these

13 patients had normal preoperative ventriculograms and were not part of this study. From the remaining 37, 32 were men and 5 women; their ages ranged from 36 to 65 years; the mean for both men and women being 50 years.

Selective coronary arteriography and bypass graft opacification were obtained with preshaped polyethylene catheters introduced by a percutaneous trans-femoral route, a method described previously.⁹ Cineventriculography was also performed through a retrograde percutaneous femoral approach.

The left ventricle was considered normal when synchronous concentric and uniform contraction of all its segments could be demonstrated. In contrast to the classification of other investigators,¹⁰ we have limited the term asynergy to the following two abnormalities of contraction: (1) akinesis indicating absence of wall movement and (2) hypokinesis indicating incomplete wall movement.

Cases showing paradoxical movement

From the Departments of Medicine and Radiology, Montreal Heart Institute, Montreal, Canada.
Supported by the Jean-Louis Lévesque Foundation.
Received for publication November 16, 1972.
Reprint requests to Dr. Michel Chabot, 5000 Belanger St. East, Montreal, Canada.

Table V Outcome of LV asynergy after operation

| Asynergy | ECG-VCG | | | | | | | | | | | | Total no |
|----------------------------|---------------|---|---|---------------|---|---|------|---|---|-----------|---|----|----------|
| | Inferior wall | | | Anterior wall | | | Apex | | | All sites | | | |
| | U* | I | N | U | I | N | U | I | N | U | I | V | |
| <i>Hypokinesis</i> | | | | | | | | | | | | | |
| Without infarction pattern | 0 | 0 | 2 | 2 | 1 | 4 | 3 | 2 | 9 | 5 | 3 | 15 | 23 |
| With infarction pattern | 0 | 0 | 0 | 3 | 1 | 1 | 2 | 3 | 0 | 5 | 4 | 1 | 10 |
| <i>Akinesis</i> | | | | | | | | | | | | | |
| Without infarction pattern | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 2 |
| With infarction pattern | 7 | 0 | 0 | 2 | 0 | 0 | 4 | 0 | 0 | 13 | 0 | 0 | 13 |

*U = horizontal or sagging ST depression; N = normal contraction.

whereas hypokinesis with infarction improved completely in only one of 10 cases ($p < 0.01$). The site of asynergy whether anterior or apical did not matter. Of the anterior wall hypokinesis seven of 12 instances improved (58.3 per cent) whereas 14 of 19 cases of apical hypokinesis also showed a better contraction (73.6 per cent). Only two instances of inferior wall hypokinesis were observed and both returned to normal contraction. Figs 1 and 2 illustrate the fate of left ventricular asynergy in a case of inferior wall akinesis and hypokinesis of the anterior and apical segments. The inferior wall akinesis persisted after operation in this patient with an inferior wall infarction whereas the contraction of the anterior and apical segments returned to normal. As shown in Table III no relationship between ischemic changes identified by horizontal or sagging ST depression with or without negative T wave and outcome of left ventricular wall asynergy could be observed since such alterations were rarely found without evidence of infarction.

The following observations may be stressed: (1) hypokinesis improved frequently whereas akinesis remained unchanged; (2) hypokinesis without evidence of old myocardial infarction was frequently completely corrected following operation whereas hypokinetic areas associated with an old myocardial infarction seldom return to normal contraction.

Discussion

Left ventricular asynergy may be due to scar tissue of previous myocardial infarction or to myocardial ischemia. From our observations it appears that akinesis is most frequently related to an extensive myocardial infarction and hence almost always irreversible. The only two instances of akinesis without evidence of infarction were located at the inferior wall where it may have been undetected because of the known limitation of these techniques in the diagnosis of infarction in that particular area. Akinesis observed in this series never improved in spite of correction of the circulatory deficit by aorta to coronary bypass graft and was most likely due to noncontractile scar tissue of previous infarction.

Hypokinesis however may or may not be associated with myocardial infarction. The improvement following this type of surgery of most hypokinetic walls without evidence of previous myocardial infarction indicates that a circulatory deficit was responsible and that these asynergic walls contained viable ischemic muscle which may contract better after operation. The lack of improvement may be explained in three of the five instances of hypokinesis without infarction. Unimproved hypokinesis of the anterior and apical walls was observed in a patient with an aorta to anterior descending graft which had a stenosis reducing its lumen by 50 per cent possibly responsible for persisting ischemic

Table II Site of asynergy and of myocardial infarction as diagnosed by ECG VCG (excluding LVH and LBBB)

| Site of asynergy | Site of MI | | | | | |
|------------------|------------|---------------|--------------|--------------|----------|----------------|
| | Anterior | Anterolateral | High lateral | Anteroseptal | Inferior | True posterior |
| ECG | | | | | | |
| Anterior | — | 2 | — | 3 | — | — |
| Apex | — | 3 | — | 7 | — | — |
| Inferior | — | — | — | — | 7 | 1 |
| VCG | | | | | | |
| Anterior | — | 2 | — | 4 | — | — |
| Apex | — | 3 | — | 7 | — | — |
| Inferior | — | — | — | — | 7 | 1 |

Table III Distribution of asynergy and ECG findings (excluding MI LVH and LBBB)

| Site of asynergy | ECG abnormalities | | |
|------------------|-------------------|--------------------------|------------|
| | Ischemic changes | Nonspecific ST T changes | Normal ECG |
| Anterior | — | 3 | 3 |
| Apex | 3 | 5 | 7 |
| Inferior | 1 | 1 | 2 |

Table IV Correlation between ECG evidence of old MI and type of asynergy with respect to the site of asynergy and localization of ECG abnormalities

| ECG VCG evidence of infarction | No | Type | |
|--------------------------------|----|----------|-------------|
| | | Ikinesis | Hypokinesis |
| Inferior and true posterior | 7 | 7 | 0 |
| Anterior* wall and/or apex | 16 | 6 | 10 |

*Including anterolateral high lateral and anteroseptal

lesions of the left main trunk, left anterior descending, and its first diagonal branch or left circumflex artery. Infarctions of the anterior and true posterior areas were re-

lated to lesions of the right coronary artery (if right coronary dominant) or left circumflex (if left circumflex dominant).

Table III relates the distribution of asynergy and ECG findings other than MI and excluding LVH and LBBB.

All investigation procedures mentioned above were performed between two and three weeks prior to operation and reported about three weeks following the operation.

Results

A good correlation exists between LCG evidence of old MI and abnormal ventricular contraction with respect to the site of asynergy on the ventriculogram and localization of the infarction pattern by ECG VCG (Table IV).

As shown in Table V the outcome of asynergy of ventricular walls inferior, anterior and apical is described for hypokinesis and akinesis in both conditions with and without ECG evidence of old myocardial infarction. It may be noted that 23 of the 33 instances of hypokinesis (70 per cent) improved whereas akinesis remained unchanged. Furthermore 19 of 23 instances of hypokinesis without LCG evidence of infarction improved (75.2 per cent) whereas only five of 10 instances of hypokinesis with infarction showed a similar favorable outcome ($p = NS$). It is also of interest to note that 15 of the 23 cases of hypokinesis without ECG pattern of infarction returned to normal contraction (65 per cent).

whereas evidence of an old myocardial infarction may predict a partial improvement only

Summary

The preoperative ECGs and VCGs of 37 patients were studied to determine whether these techniques might help to forecast the outcome of left ventricular asynergy following aortocoronary saphenous vein graft surgery. It was found that akinesis was always associated with a pattern of old myocardial infarction and that it always remained unchanged after operation. Hypokinesis however was frequently improved. The incidence and degree of improvement were somewhat related to the presence or absence of old myocardial infarction. Hypokinesis without myocardial infarction improved lightly more frequently than that associated with a myocardial infarction (78.2 vs 50 per cent). However 60 per cent of the cases of hypokinesis without myocardial infarction returned to normal contraction following operation whereas only 10 per cent of the cases with hypokinesis and myocardial infarction were completely improved. These findings suggest that the preoperative ECG and VCG may reasonably predict the outcome of hypokinesis following aortocoronary bypass surgery, but only in terms of complete recovery which is most frequently observed in hypokinesis without ECG evidence of old myocardial infarction.

REFERENCES

- 1 Favaloro R G Saphenous vein graft in the surgical treatment of coronary artery disease *J Thorac Cardiovasc Surg* 58:178 1969
- 2 Eifler D B Favaloro R G and Groves L K Coronary artery surgery utilizing saphenous vein graft techniques Clinical experience with 224 operations *J Thorac Cardiovasc Surg* 59:147 1970
- 3 Mitchell B F Adam M Lambert C J Singu U and Sheikh S Ascending aorta to coronary artery saphenous vein bypass grafts *J Thorac Cardiovasc Surg* 60:457 1970
- 4 Johnson W D Fleimms R J and Lepley D Jr Direct coronary surgery utilizing multiple vein bypass grafts *Ann Thorac Surg* 9:436 1970
- 5 Field I Trimble A S Trobridge G F and Aldridge H E Direct myocardial revascularization on saphenous vein bypass grafts to the distal coronary artery *Ann Thorac Surg* 10:112 1970
- 6 Bigelow W G Surgical treatment of coronary heart disease *Can Med Assoc J* 101:404 1971
- 7 Saltiel J Lesperance J Bourassa M G Castonguay Y Campeau L and Grondin P Reversibility of left ventricular dysfunction following aortocoronary bypass grafts *Am J Roentgenol Radium Ther Nucl Med* 117:139 1970
- 8 Bourassa M G Lesperance J Campeau L and Saltiel J Fate of left ventricular contraction following aortocoronary venous grafts Early and postoperative modifications *Circulation* 46:724 1972
- 9 Bourassa M G Lesperance J and Campeau L Selective coronary arteriography by the percutaneous femoral artery approach *Am J Roentgenol Radium Ther Nucl Med* 107:1377 1969
- 10 Herman M V Heintz R A Klein M D and Gorlin R Localized disorders in myocardial contraction *N Engl J Med* 277:722 1967
- 11 Masse E and Walsh T J Clinical vector cardiography and electrocardiography Chicago 1960 Year Book Medical Publishers Inc
- 12 Gunnar R M Pietras R J Blackaller J Damm S E Szanto P B and Tobin J R Jr Correlation of vectorcardiographic criteria for myocardial infarction with autopsy findings *Circulation* 30:158 1967
- 13 Myers G B Klein R A and Hiratzka T Correlation of electrocardiographic and pathological findings in posterior infarction *AM HEART J* 38:547 1949
- 14 Okamoto N Simonson E Abuys S and Vanning C Significance of the initial R wave in lead aVR of the electrocardiogram in the diagnosis of myocardial infarction *Circulation* 30:176 1967
- 15 Perloff J K The recognition of strictly posterior myocardial infarction by conventional scalar electrocardiography *Circulation* 30:106 1964
- 16 Lickissen J EKG in strictly posterior myocardial infarction *Acta Med Scand* 187:465 1970
- 17 Doucet P Walsh T J and Masse E A vectorcardiographic and electrocardiographic study of left bundle branch block with myocardial infarction *Am J Cardiol* 14:171 1966
- 18 Chou T and Helm R A Clinical vector cardiography New York 1967 Grune & Stratton Inc
- 19 Young E and Williams C The frontal vector cardiogram in old inferior myocardial infarction *Circulation* 37:604 1968
- 20 Mayhew V S and Levine H D Vectorcardiographic differentiation between right ventricular hypertrophy and posterobasal myocardial infarction *Circulation* 42:883 1970
- 21 McConahay D R McCallister B D Hallerman F J and Smith R E Comparative quantitative analysis of the electrocardiogram and the vectorcardiogram Correlations with the coronary arteriogram *Circulation* 42:245 1970



Fig 2 A and B The opacified left ventricle in end diastole (A left) and in end systole (B right) after operation

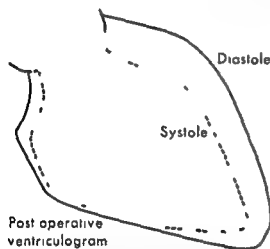


Fig 2C Comparison of these images shows a return to normal contraction of the anterior and apical segments but a persisting akinesis of the inferior wall

changes. In another patient with unimproved hypokinesis of the anterior and apical segments, ECG changes during the early postoperative course suggested an anterior wall infarction. As for the only instance which remained unexplained one may assume that chronic ischemia of long standing may have produced irreversible muscle changes other than a circumscribed myocardial infarction or that the myocardial infarction was not detected. The partial improvement of the three instances of hypokinesis without infarction may be explained on the same basis.

On the other hand, hypokinesis associated with an ECG pattern of old myocardial infarction seldom returns to normal contraction in spite of an apparent adequate correction of the circulatory deficit sug-

gesting that scar tissue is the most significant factor. In fact, only one such case showed a return to normal contraction. It should be noted, however, that the infarction was evident only on the VCG, suggesting a small area of fibrosis which may not have altered the ventricular contraction pattern as observed at ventriculography. In cases where a partial recovery was observed it is likely that previously ischemic but viable muscle intermingled with or surrounding fibrous tissue became better perfused following operation and hence showed a better contraction.

Our study confirms the reports of others which have indicated a fairly good correlation between zones of altered ventricular contraction and sites of coronary artery narrowing.²¹ It has also demonstrated that infarctions as documented by ECG and VCG are appropriately located in relation to ventricular zones of asynergy.

Our results suggest that the preoperative LCG VCG in patients may predict the outcome of left ventricular contraction following aortocoronary bypass grafts only to a limited degree. First the presence or absence of ECG evidence of an old myocardial infarction is the only helpful criterion. Second this criterion may be helpful particularly in predicting the outcome of hypokinesis. In fact akinesis rarely improves at least in our experience and it is almost invariably associated with an old myocardial infarction. As far as hypokinesis is concerned the absence of LCG pattern of an old myocardial infarction may forecast a return to normal contraction.

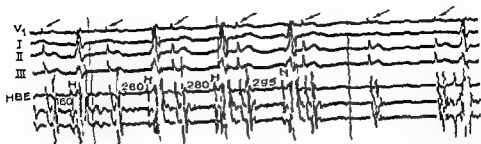


Fig 1 Electrophysiological recordings from Case 1. Shown are Leads V₁, I, II and III with His bundle electrogram (HBE). Pacing impulses are labeled with arrows. His bundle electrograms are labeled H. The pacing rate is 120 per minute and P-H interval are fixed and show Wenckebach periodicity. The fifth and sixth P waves are blocked proximal to the H potential. Paper speed is 100 mm per second and time lines are at 1 second.

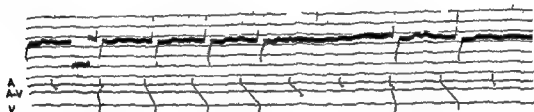


Fig 2 Rhythm strip and ladder diagram showing Wenckebach period terminated with two blocked P waves. The upper level of the ladder reflects atrium, the middle two levels A-V node, and the bottom ventricle. The third, fourth, and fifth P waves are conducted with increasing P-R intervals. The sixth and seventh P waves are blocked. A proposed mechanism is presented showing the first P blocked low in the node and the second P blocked at a higher level. (From Langendorf and Pick, *Circulation* 13:381, 1956, reproduced with permission of the American Heart Association.)

0.10 second and a P-R interval of 0.14 second. Incomplete right bundle branch block pattern was also noted.

Electrophysiological studies. Informed consent was obtained for electrophysiological studies. His bundle electrograms were recorded with the use of previously described techniques.¹¹ Atrial pacing was performed at varied heart rates.

The patient was in sinus rhythm at a rate of 60 per minute. P-H interval was 125 msec, P-A 20 msec, A-H 105 msec, and H-V 40 msec. Wenckebach periods proximal to the His bundle were noted at a paced rate of 120 per minute. These were generally terminated by one blocked P. Several atypical cycles were recorded, these being terminated by two consecutive P waves blocked proximal to the His bundle (Fig 1).

Comment. Langendorf and Pick² postulated that block of two consecutive P waves during a Wenckebach period occurs due to inhomogeneous penetration of the A-V node. They suggested deeper penetration of the first impulse and more superficial penetration of the second impulse (Fig 2). His bundle electrograms in such cases should record block of both P waves proximal to the His bundle.

In the present case His bundle electro-

grams documented repetitive block proximal to the His bundle. This technique does not allow us to measure the degree of penetration of the A-V node in the two blocked cycles. However, the results are consistent with Langendorf and Pick's previous hypothesis.

Case 2. A 25 year old man with corrected tetralogy of Fallot was admitted for cardiac evaluation. ECG revealed normal sinus rhythm at a rate of 70 per minute, a P-R interval of 0.16 second, and a QRS duration of 0.16 second with complete right bundle branch block pattern.

Electrophysiological studies. The patient was in sinus rhythm at a rate of 65 per minute. The conduction intervals were as follows: I-H 178 msec, P-A 23 msec, A-H 105 msec, and H-V 44 msec. 4:3 and 3:2 Wenckebach periods proximal to H were noted at a paced rate of 150 per minute. The Wenckebach periods were typical during 4:3 conduction (Fig 3 A). With 3:2 A-V nodal Wenckebach the following was observed. The first P-H was 140 msec, the second 160 msec, and the third P was blocked proximal to the His bundle. The second P was always blocked distal to H (Fig 3 B).

Comment. In this case 3:2 Wenckebach periods proximal to H were associated with 3:1 A-V block. Inhomogeneous conduction

Wenckebach periods with repetitive block

Evaluation with His bundle recording

Ramesh C Dhinra MD

Kenneth M Rosen, MD, FACC

Shahbudin H Rahimtoola, FRCP

Chicago, Ill

Langendorf¹ in 1948 introduced the term "concealed conduction" to describe the results of partial penetration of an electrical impulse into the A V junction. The incompletely penetrating impulse transiently lengthens the A V refractory period modifying the conduction of the subsequent impulse. This concept has proved useful in understanding the mechanisms involved in a number of arrhythmias.^{2,3} Repetitive block of several consecutive atrial impulses can occur when successive impulses partially penetrate the A V conducting system without traversing it completely, producing varying periods of ventricular asystole. This phenomenon has been called "repetitive concealed conduction."²

Type I second degree A V block (Wenckebach) is characterized by progressive PR prolongation prior to a single dropped beat. Termination of a Wenckebach cycle with more than one blocked P wave is rare. Langendorf and Pick² reported a patient with Wenckebach periods terminated by several blocked atrial impulses and suggested that this was a mani-

festation of repetitive concealed conduction.

In this report we describe two patients with Wenckebach periods and block of two consecutive P waves. In one the mechanism appeared similar to that previously postulated by Langendorf and Pick. In the second a new mechanism is described. In this case long cycle lengths were produced in the His Purkinje system due to A V nodal Wenckebach periods. This long cycle prolonged the His Purkinje refractory period in the subsequent cycle so that the second conducted beat of the Wenckebach period (short cycle) was blocked distal to the His bundle. This occurred during 3:1 A V nodal Type I block with resulting 3:1 A V block.

Both mechanisms can produce unexpected asystole during Type I second degree A V block.

Report of cases

Case 1 A 23 year old man was admitted to Cook County Hospital for evaluation of syncope. Results of a physical examination were essentially negative. Electrocardiogram (ECG) revealed sinus bradycardia at a rate of 53 per minute, a QRS duration of

From the Department of Adult Cardiology, Cook County Hospital, and the Section of Cardiology, Department of Medicine, The Abraham Lincoln School of Medicine, University of Illinois College of Medicine, Chicago, Ill.
Supported in part by NIH contract 71-2478 under the Myocardial Infarction Program, National Institutes of Health, Department of Health, Education and Welfare.
Received for publication Nov. 16, 1972.
Reprint requests to Kenneth M. Rosen, MD, Section of Cardiology, Department of Medicine, University of Illinois Hospital, 840 South Wood St., Chicago, Ill. 60612.

attributed this to repetitive concealed conduction in the A V junction suggesting that the first blocked sinus impulse had a deeper penetration of the A V node augmenting its refractoriness so that the subsequent sinus impulse could not be transmitted. It was postulated that this latter impulse was blocked at a higher level. Such inhomogeneous penetration of the A V junction could be responsible for block of multiple atrial impulses.

Moore and associates¹⁰ experimentally demonstrated repetitive concealed conduction with as many as six premature atrial impulses blocked in succession. Watanabe and Dreifus¹¹ demonstrated two levels of A V nodal block during 3:1 response in rabbit hearts and suggested inhomogeneous conduction in the A V node. Moore¹² employing microelectrode technique in rabbits defined the sites and mechanisms of A V block of serially concealed premature atrial impulses and demonstrated decremental conduction within both the A V node and His Purkinje system.

His bundle recording has helped clarify mechanisms of concealed conduction in man, permitting accurate localization of sites of conduction delay following blocked premature responses.^{13,14} This technique was helpful in the present cases. In the first patient repetitive block during Wenckebach beating was localized proximal to the His bundle. This is in keeping with the postulation of inhomogeneous conduction in the A V node as previously suggested.¹

In the second patient varying depths of penetration were documented with His bundle recording. Block of two successive P waves occurred, one distal and the other proximal to the His bundle. The consecutive blocked P waves did not reflect repetitive concealed conduction. Failure of impulse propagation at the A V node (during Type I block) produced long cycle lengths in the His Purkinje system with associated refractory period prolongation. This allowed the second beat of the Wenckebach period (short cycle) to be blocked distal to His bundle. Thus 3:2 A V nodal Wenckebach periods were associated with 3:1 A V block. It should be pointed out that the initiation of this 3:1 block must have oc-

curred with a typical 3:2 Wenckebach period.

Clinical implications. Although the abnormalities of conduction demonstrated were induced by atrial pacing, similar phenomenon could certainly occur during spontaneous beating explaining unexpected blocked P waves during Type I second-degree block. In patients with bundle branch block, long R-R cycles could lengthen the refractory period of the functioning bundle branch, causing nontransmission during short subsequent cycles. 3:1 A V block with bundle branch block could thus represent one of the following: (1) 3:1 block proximal to H, (2) 3:1 block distal to H, or (3) the mechanism described in Case 2. Mechanism No. 2 would appear to be the most serious, suggesting bilateral bundle branch disease.¹⁵

Both types of repetitive block described in this report could produce serious bradyarrhythmias during the course of usually benign Type I second-degree A V block. Pharmacologic therapy with atropine or isoproterenol or pacemaker insertion might be necessary.

Summary

Two patients are reported in whom repetitive block of two consecutive P waves occurred during Wenckebach beating induced by atrial pacing. His bundle recordings revealed block proximal to H in the first case suggesting inhomogeneous conduction in the A V node. In the second case long cycle lengths were produced in the His Purkinje system due to A V nodal Wenckebach periods. The long cycles prolonged refractory periods in the His Purkinje system so that subsequent beats (short cycles) were blocked distal to H.

The repetitive block of consecutive multiple atrial impulses could result in unexpected degrees of ventricular asystole during usually benign Type I second-degree A V block.

The authors are grateful to Drs. Richard Langendorf and Alfred Pick for their kind permission to use Fig. 2 from their previous publication. The authors also wish to express their thanks to Mrs. Mary Ellen Rosen for the preparation of the illustrations and to Mrs. Katherine Evans for the preparation of the manuscript.



Fig 3 Wenckebach periods produced with atrial pacing demonstrating block above and below the His bundle. The pacing rate is 150 per minute in both panels. (A) 4:3 Wenckebach periods proximal to the His bundle. Note the long H-H cycle of 730 msec and the short H-H cycle of 435 msec. (B) 3:2 A-V nodal Wenckebach with 3:1 block. The second P wave of each cycle is blocked distal to H and the third proximal to H. The long H-H cycles are 790 msec and the short 435 msec. (See text for discussion.) Paper speed is 100 mm per second and the time lines are at 1 second.

was noted the first P being blocked distal to and the second proximal to the His bundle.

The failure of conduction distal to H of the first nonpropagated impulse during 3:1 response was easily explainable. It is evident from Fig 3 B that H potential occurred at alternating long and short intervals. The longer H-H cycles were of 790 msec duration while the shorter cycles were 435 msec. This long-short sequence reflected 3:2 Wenckebach periodicity proximal to H. The refractory period of the ventricular specialized conduction system was presumably prolonged in the cycle following the long H-H, reflecting the relationship of cycle length and refractory period in the His-Purkinje system.⁸ This prolongation of refractoriness was reflected in block distal to H of the second beat of the Wenckebach sequence (short H-H cycle). Since the patient had right bundle

branch block block distal to H probably occurred in the left bundle branch.

The above interpretation was supported by examining conduction during the 4:3 Wenckebach periods. During 4:3 conduction the short H-H cycles were 435 msec identical in length to short cycles during 3:2 conduction. The long H-H cycles were only 730 msec and shorter than the cycles during 3:1 conduction. This slight shortening of long H-H cycles was of critical degree allowing conduction to occur in the subsequent short cycle.

Discussion

In 1925 Lewis and Master⁹ demonstrated that a blocked atrial impulse could delay the transmission or completely block a subsequent impulse. They also described block of repetitive impulses. Langendorf and Pick² described block of consecutive impulses during a Wenckebach cycle. They

Electrocardiographic findings in single ventricle and related conditions

M Quera Jimenez MD
M Casanova Gomez MD
C Castro Gussons MD
F Moreno-Granado MD
V Perez Martinez MD
G Merino Baltes MD
Madrid Spain

Although this subject has been studied for a relatively long time the electrocardiographic features reported by different authors have become more specific as knowledge in the pathology of these malformations has increased.¹⁻³ It is therefore desirable that analysis be based on groups with maximal anatomic and physiological homogeneity. Thus two main groups have been established (I and II) with a further three subgroups each (A, B and C) (See Table I).

In Group I (Table I) both atrioventricular orifices either independent or fused into a common atrioventricular canal are mainly received by the left ventricle. In this group where a total or partial lack of the normal displacement of the atrioventricular canal towards the bulbus cordis appears to be the basic embryologic derangement.⁴

Defective displacement of the atrioventricular canal towards the bulbus cordis would be the adequate term. Subgroups A, B and C concern the extent with which this normal displacement has been lacking.

In Subgroup A (Table I) the atrioventricular canal is completely received by the left ventricle. The result is a bigger left ventricular cavity in comparison with the right ventricle which is reduced to its outflow tract. This has previously been described as single left ventricle,⁵ single (primitive) ventricle,⁶ double inlet left ventricle,⁷ displaced tricuspid valve,⁸ etc. Common associated malformations with this anomaly are transposition of the great arteries, pulmonary stenosis,⁹ preductal coarctation of the aorta, subaortic stenosis, etc. Less frequent are certain anomalies of the atrioventricular valves including mitral atresia.¹⁰

In Subgroup B (Table I) the normal displacement of the atrioventricular canal towards the bulbus cordis, although initiated, would remain somewhat incomplete. Consequently despite a small area of the atrioventricular canal being received by the right ventricle, the major portion of this connects with the left ventricular cavity. The resulting ventricular proportions, although very similar to those in Subgroup A,

From the Servicio de Cardiología Pediátrica de la Clínica Infantil de la Seguridad Social, La Paz, Madrid, Spain.
Received for publication May 20, 1972.

Reprints: Dr. M. Quera Jimenez, Jefe del Servicio de Cardiología Pediátrica de la Clínica Infantil de la Paz, Madrid, Spain.

REFERENCES

- 1 Langendorf R Concealed A V conduction The effect of blocked impulses on the formation and conduction of subsequent impulses *Am Heart J* 35:542 1948
- 2 Langendorf R and Pick A Concealed conduction Further evaluation of a fundamental aspect of propagation of the cardiac impulse *Circulation* 13:381 1956
- 3 Langendorf R Pick A Edelist A and Katz I N Experimental demonstration of concealed A V conduction in the human heart *Circulation* 32:386 1965
- 4 Scheiner L B and Stock R J Coupled pacing and coupled pacing with concealed conduction Report of a case describing a new observation *Circulation* 34:759 1966
- 5 Carleton R A and Graettinger J S Evidence of concealed atrioventricular conduction in man *Circulation* 31:756 1966
- 6 Scherlag B J Lau S H Helfant R H Berkowitz W D Stein E and Damato A N Catheter technique for recording His bundle activity in man *Circulation* 39:13 1969
- 7 Narula O S Cohen L S Samet P Lister J W Scherlag B and Hildner F J Localization of A V conduction defects in man by recording of the His bundle electrogram *Am J Cardiol* 25:228 1970
- 8 Moe G K Mendez C and Han J Aberrant A V impulse propagation in the dog heart A study of functional bundle branch block *Circ Res* 16 261 1965
- 9 Lewis T and Master A M Observations upon conduction in the mammalian heart A V conduction *Heart* 12 209 1925
- 10 Moe G K Abildskov J A and Mendez C An experimental study of concealed conduction *Am Heart J* 67:338 1964
- 11 Watanabe Y and Dreifus L S Second degree atrioventricular block *Cardiovasc Res* 1:150 1967
- 12 Moore N Microelectrode studies in concealment of multiple premature atrial responses *Circ Res* 12 660 1966
- 13 Damato A N and Lau S H Concealed and supernormal atrioventricular conduction *Circulation* 43:967 1971
- 14 Rosen K M Rahimtoola S H and Gunnar R M Pseudo A V block secondary to premature nonpropagated His bundle depolarizations Documentation by His bundle electrocardiography *Circulation* 42:367 1970
- 15 Langendorf R and Pick A Atrioventricular block type II (Mobitz) Its nature and clinical significance *Circulation* 38 819 1968

completely received by the right ventricle which is greatly enlarged and may appear as a single ventricular chamber. The left ventricle deprived from any significant blood filling remains reduced to a very hypoplastic cavity without papillary muscles. It is questionable whether these cases are not equivalent to those described as single right ventricle² although the finding of the remaining left ventricle has not been quite clear in the latter. Establishing a parallel with the recent contribution of Liberthson and associates⁷ these cases could equally be termed as *displaced mitral valve*¹⁴.

In Subgroup B (Table I) the atrioventricular canal abnormally displaced towards the bulbus cordis remains slightly connected to the left ventricle. This cavity provided with the posterior group of papillary muscles and accessible to blood filling ■ of larger size.

In Subgroup C (Table I) the exaggerated displacement of the atrioventricular canal towards the bulbus cordis is of lesser degree. A comparatively larger area of atrioventricular canal than in Subgroups A and B ■ received by the left ventricle the cavity of which shelters the two normal groups of papillary muscles and attains a more regular size.

Special case In one case with the mitral orifice-ventricular septum relationship fairly similar to the one in Subgroup C and the left ventricle containing only the posterior group of papillary muscles as in Subgroup B we were surprised to find the left ventricle considerably larger than the right. This unusual factor is ascribed to the hypoplasia of the tricuspid orifice despite part of the mitral orifices being received by the right ventricle the total area of the atrioventricular canal connected to the left ventricle was larger.

Material and methods

Of a total of 33 cases studied an exact anatomical diagnosis was obtained in 25 with a fairly reliable diagnostic approach being made in the remainder using all the conventional methods including cardiac catheterization and angiocardiology.

The angiocardiology difference between Group 1A, 1B and 1C was based on the presence of the characteristic filling

defects (interpreted as papillary muscles) in the remainder of the right ventricular cavity. Angiocardiology differentiation between groups 1A and 1B is not considered feasible.

Apart from the basic pertaining sub-group other associations (type of bulbo-ventricular loop², pulmonary stenosis and common atrioventricular canal) have been taken into account in analyzing the electrocardiograms of the cases concerned.

For general case distribution incidence rates of most frequent associations see Table I.

Normal atrial and visceral position was detected in every case. A ventricular septal defect was present in all cases in Group I and in all but one case of Subgroup A in Group II.

The electrocardiographic criteria used for the diagnosis of atrial and ventricular enlargement have been those proposed by others^{15, 17}.

Results

In grouping the cases for the electrocardiographic analysis the main factors under consideration were the type of bulboventricular loop and the presence or absence of pulmonary outflow obstruction and dextrocardia.

1 Group I Subgroups A and B D loop Subgroups I A and I B have been analyzed together since there were no significant differences among them.

A NO PULMONARY STENOSIS Five cases (1 to 5 Table II). There was a sinus rhythm in every case. There was a right atrial enlargement in cases 1 to 4; in case 5 biatrial enlargement was diagnosed.

The QRS axis (Fig. 1) was of almost +100 degrees in three cases being of -80 degrees and -10 degrees in cases 3 and 4 respectively. A common atrioventricular valve and normally related great arteries were found in case 3. Deep Q waves in Leads III and aV_F were absent in all cases. Essentially negative complexes in the Lead aV_R were found in cases 1, 2, 4 and 5. Case 3 showed a qR deflection in the right arm lead. The initial QRS vector in the horizontal plane was directed towards the right anteriorly in all cases (Fig. 2).

RS complexes over the entire precordium were noted in cases 1 and 3. ST-T digitalis








| GROUP I | | | NORMAL | GROUP II | | |
|--|---|---|---|---|---|---|
| A | B | C | | C | B | A |
|  |  |  |  |  |  |  |
| 11 | 6 | 2 | TOTAL CASES 33* | 1 | 2 | 2 |
| D-LOOP 4 L-LOOP 7 | D-LOOP 2 L-LOOP 4 | D-LOOP 1 L-LOOP 1 | B-V LOOP | D-LOOP 1 | D-LOOP 2 | D-LOOP 2 |
| 2A-VV 8 CA-VC 2 MA 1 | 2A-VV 4 CA-VC 1 MA 1 | 2A-VV 1 CA-VC 1 | AV VALVES | 2A-VV 1 | CA-VC 2 | 2A-VV 1 CA-VC 1 |
| 4 | 1 | 1 | PULMONARY ATRESIA/STENOSIS | — | 1 | — |
| L 10 R 1 | L 6 R — | L 2 R — | CARDIAC APEX | L | L | L |

Table 1 General distribution and classification of our cases. The special case—the 3 dextrocardia cases and the angiocardigraphically proved cases have not been included in this table. See text for criteria used to classify the cases. RV = right ventricle; LV = left ventricle; TPM = tricuspid papillary muscle; MPM = mitral papillary muscle; AVV = atrioventricular valve; CAVC = common atrioventricular canal; R = right; L = left; D = dextro; L = levo.

will vary somewhat from the right ventricular size. Effectively the right ventricle receiving a small but definite part of the atrioventricular canal has a true but small sinusal portion with papillary muscles and chordae. The cases included in Subgroup II generally come under the same designations as used for those in Subgroup A; the associated anomalies being similar. Nevertheless, we have established two separate groups to ascertain whether such a small difference in the size of the right ventricle is reflectable in the electrocardiogram.

In Subgroup C (Table 1) the displacement of the atrioventricular canal towards the bulbus cordis appears nearer to the normal. Consequently, a larger area of the atrioventricular canal is received by the right ventricle, the size of which would significantly differ from that in Subgroups A and B. The terms of Lambert heart and

straddling tricuspid valve^{7,10,11} have been used so far to designate malformations with these general characteristics. One case with an associated mitral atresia has also been reported in this subgroup.¹²

Less common and in turn lesser known are the hearts studied in Group II.^{2,3,6} As for the atrioventricular canal-ventricular septum relationships the opposite phenomenon characterizing the cases of Group I seems to be the basic feature of the ones in Group II. Thus the term "Exaggerated displacement of the atrioventricular canal towards the bulbus cordis" is indicated for these malformations.^{13,14}

As in Group I three further subgroups, A, B, and C, have been established in accordance with the extent of this exaggerated displacement. In Subgroup A (Table 1) the atrioventricular canal common or divided in two atrioventricular orifices, is

completely received by the right ventricle which is greatly enlarged and may appear as a single ventricular chamber. The left ventricle deprived from any significant blood filling remains reduced to a very hypoplastic cavity without papillary muscles. It is questionable whether these cases are not equivalent to those described as single right ventricle² although the finding of the remaining left ventricle has not been quite clear in the latter. Establishing a parallel with the recent contribution of Libberthson and associates³ these cases could equally be termed as displaced mitral valve.¹⁴

In Subgroup B (Table I) the atrioventricular canal abnormally displaced towards the bulbus cordis remains slightly connected to the left ventricle. This cavity provided with the posterior group of papillary muscles and accessible to blood filling is of larger size.

In Subgroup C (Table I) the exaggerated displacement of the atrioventricular canal towards the bulbus cordis is of lesser degree. A comparatively larger area of atrioventricular canal than in Subgroups A and B is received by the left ventricle; the cavity of which shelters the two normal groups of papillary muscles and attains a more regular size.

Special case. In one case with the mitral orifice-ventricular septum relationship fairly similar to the one in Subgroup C and the left ventricle containing only the posterior group of papillary muscles as in Subgroup B we were surprised to find the left ventricle considerably larger than the right. This unusual factor is ascribed to the hypoplasia of the tricuspid orifice despite part of the mitral orifices being received by the right ventricle; the total area of the atrioventricular canal connected to the left ventricle was larger.

Material and methods

Of a total of 33 cases studied an exact anatomical diagnosis was obtained in 23 with a fairly reliable diagnostic approach being made in the remainder using all the conventional methods including cardiac catheterization and angiocardiology.

The angiocardiological difference between Group IA, IB and IC was based on the presence of the characteristic filling

defects (interpreted as papillary muscles) in the remainder of the right ventricular cavity. Angiocardiological differentiation between groups IA and IB is not considered feasible.

Apart from the basic pertaining subgroup other associations (type of bulboventricular loop² pulmonary stenosis and common atrioventricular canal) have been taken into account in analyzing the electrocardiograms of the cases concerned.

For general case distribution incidence rates of most frequent associations see Table I.

Normal atrial and visceral position was detected in every case. A ventricular septal defect was present in all cases in Group I and in all but one case of Subgroup A in Group II.

The electrocardiographic criteria used for the diagnosis of atrial and ventricular enlargement have been those proposed by others.^{15, 17}

Results

In grouping the cases for the electrocardiographic analysis the main factors under consideration were the type of bulboventricular loop and the presence or absence of pulmonary outflow obstruction and dextrocardia.

I Group I Subgroups A and B D loop. Subgroups IA and IB have been analyzed together since there were no significant differences among them.

A NO PULMONARY STENOSIS. Five cases (1 to 5 Table II). There was a sinus rhythm in every case. There was a right atrial enlargement in cases 1 to 4; in case 5 biatrial enlargement was diagnosed.

The QRS axis (Fig. 1) was of almost +100 degrees in three cases, being of -80 degrees and -10 degrees in cases 3 and 4 respectively. A common atrioventricular valve and normally related great arteries were found in case 3. Deep Q waves in Leads III and aV₁ were absent in all cases. Essentially negative complexes in the Lead aV₂ were found in cases 1, 2, 4 and 5. Case 3 showed a qR deflection in the right arm lead. The initial QRS vector in the horizontal plane was directed towards the right anteriorly in all cases (Fig. 2).

RS complexes over the entire precordium were noted in cases 1 and 3. ST-T digitalis

Table II Groups IA and IB—"D" bulboventricular loop—ECG findings

| Case No and initials | Sub group | Age | Diagnosis | A V values | GA* | Rhythm |
|----------------------|-----------|-----------------|-----------|------------|--------|--------|
| Without PS | | | | | | |
| 1 R A S | A | 15 months | N | 2 | D TGA | S |
| 2 T G P | A | 15 days | N | 2 | D TGA | S |
| 3 R A P | B | 1 day/2 months | N | CAVC | Normal | S |
| 4 G G | B | 3 months | N | 2 | D TGA | S |
| 5 J M P | † | 2 years/6 years | A | ? | D TGA | S |
| With PS | | | | | | |
| 6 M S | A | 3 months | N | MA | Normal | AVC |
| 7 P F | A | 3 months | N | CAVC | D TGA | S |
| 8 A G C | ? | 1 year | A | ? | Normal | S |

*Abbreviations GA = great arteries; N = necropsy; A = angiography; CAVC = common atrioventricular canal; MA = mitral atresia; RVH = right ventricular hypertrophy; LVH = left ventricular hypertrophy; BVH = biventricular hypertrophy; PS = pulmonary stenosis; †The angiocardigraphic differentiation between group IA and IB and the prediction of the common or divided nature of the AV canal.

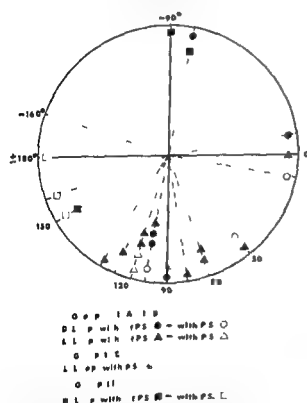


Fig 1 General distribution of the QRS axis in the frontal plane in our cases. Cases 22 and 23 are not included because of the undetermined nature of their QRS axis. The cases with dextrocardia are excluded from this illustration. Note the majority of the cases showing a QRS axis varying between $+50$ and $+130$ degrees. The three cases with an electrical axis close to -90 degrees having a common atrioventricular valve. Nevertheless, four other cases with a common atrioventricular orifice had an electrical axis varying from $+10$ to $+180$ degrees. Note in cases of Group II (with a larger right ventricle) the QRS axis is located more to the right, as is also the electrical axis of Group I cases with an L Loop.

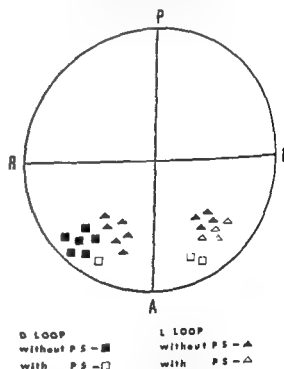


Fig 2 Group I General distribution of the initial QRS forces in the horizontal plane. Note the high incidence of abnormally directed initial vectors in the L Loop and pulmonary stenosis cases. P = posterior; I = anterior; R = right; L = left.

induced changes in the left precordial leads were present in cases 1, 3, and 4.

Electrocardiographic diagnosis of left ventricular hypertrophy was possible in cases 2, 4, and 5. In cases 1 and 3, an electrocardiogram suggesting biventricular hypertrophy was found.

B PULMONARY STENOSIS Three cases

| Atrial enlargement | A QRS | Deep Q III-aV _F | aV _R | Isodisphasicism | ST T changes | Ventricular pattern |
|--------------------|-------|----------------------------|-----------------|-----------------|--------------|---------------------|
| RAH | +100° | - | rS | + | + | BVH |
| RAH | +100 | - | rS | - | - | LVH |
| RAH | -80 | - | qR | + | + | BVH |
| RAH | -10 | - | Qr | - | + | LVH |
| BAH | +90 | - | Qr | - | - | LVH |
| RAH | +50 | - | QS | rS | + | LVH |
| - | +10 | - | QS | - | - | LVH |
| RAH | +100 | + | Qr | rS | - | LVH |

al AAHC = accelerated A to C due to RAH = right atrial hypertrophy LAH = left atrial hypertrophy BAH = bilateral hypertrophy
no considerable impossible



Fig 3 Electrocardiogram of Case 11 Complete heart block recorded in Lead I (age = 4 months)

(6 to 8 Table II) A sinus rhythm was present in all but case 6 where a short PR was found

A right atrial hypertrophy existed in cases 6 and 8. The QRS axis showed a great variability ranging from +10 to +100 degrees (Fig 1). Deep Q waves in Leads III and aV_F were noted in case 8. Essentially negative deflections were seen in Lead aV_R in every case. The initial QRS forces in the horizontal plane were abnormally directed towards the left anteriorly in two cases being normal in case 7 (Fig 2).

There was an rS deflection in all precordial leads of cases 6 and 8 and in case 6 ST T changes not induced by digitalis intake existed in the left precordial leads.

Left ventricular hypertrophy was diagnosed in every case.

II Group I Subgroup A and B L loop

Aortic aortic stenosis. Ten cases (9 to 18 Table III). A sinus rhythm was present in all except case 11 where a complete heart block existed (Fig 3). In cases

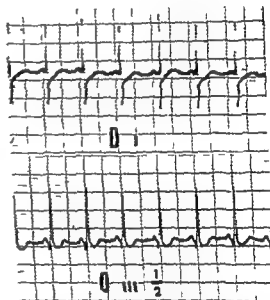


Fig 4 Electrocardiogram of Case 16. Observe the short PR segment without any other alteration in the P and QRS morphologies (accelerated atrioventricular conduction) (age = 3.5 months)

9 and 16 (Fig 4) an accelerated atrioventricular conduction was found.

Right atrial enlargement was detected in cases 10, 11, 12, 14, 15 and 17 (Fig 5) atrial hypertrophy being found in case 9. The QRS axis ranged from 0 to +120 degrees (Fig 1). Deep Q waves in Leads III and/or aV_F existed in cases 9, 11, 14, 15 and 18 (Figs 5 and 6). Essentially negative

Table III Groups IA and IB—"L" bulboventricular loop—ECG findings

| Case No and initials | Sub group | Age | Diagnosis | A V values | GA | Rhythm |
|----------------------|-----------|-----------------|-----------|------------|-------|--------|
| 9 V A R | A | 25 months | N | 2 | D TGA | AAVC |
| 10 E M L | A | 45 months | N | 2 | L TGA | S |
| 11 P P A | A | 4 months | N | 2 | L TGA | CHB* |
| 12 C A | A | 2 months | N | CAVC | L TGA | S |
| 13 P V | A | 5 months | N | 2 | L TGA | S |
| 14 L A G | B | 4 months | N | 2 | L TGA | S |
| 15 C P P | B | 3 months | N | MA | D TGA | S |
| 16 L Q D | B | 35 months | N | 2 | D TGA | AAI |
| 17 L O G | ? | 12 days/1 month | A | ? | L TGA | S |
| 18 S D | ? | 4 months | A | ? | L TGA | S |
| 19 A G | A | 1 day | N | 2 | L TGA | S |
| 20 T N T | B | 10 days | N | 2 | L TGA | S |
| 21 A P F | ? | 15 months | A | ? | L TGA | S |

*CHB = complete heart block. For other abbreviations see Table II

Table IV Group IC, ECG findings

| Case No and initials | Age | Diagnosis | A V values | GA | Rhythm | Atrial enlargement | A QRS |
|----------------------|------------------|-----------|------------|---------|--------|--------------------|-------|
| 22 S G | 1 month/3 months | N | 2 | D Malp† | S | RAH | Und |
| 23 A P J | 8 months | N | CAVC | D TGA | S | RAH | +90° |

*Abbreviations PA = pulmonary atresia Und = undetermined D Malp = D malposition For other abbreviations see Table II
†A right sided aorta and a left sided pulmonary artery arising both from a right sided morphological right ventricle. Both great arteries

deflections in Lead aV_R appeared in each case (Fig 7)

Abnormally directed initial QRS forces in the horizontal plane were noted in cases 9 13 15, and 17 a q wave in Leads V_r and V_i being detected in cases 9 and 13 (Figs 2 and 7) RS morphologies appeared in the entire precordium of cases 10 11 12 13 16, and 17 (Fig 6) Case 15 showed in rS deflection in all precordial leads (Fig 5)

Digitalis induced ST T changes over the left precordial leads were seen in cases 9 10, 12, 13, 15 16 and 18 (Figs 5 6 and 7)

Diagnosis of left ventricular hypertrophy was possible in cases 12 11 15 and 18 (Fig 8), a right ventricular hypertrophy being detected in cases 9 and 11, with biventricular enlargement in the remaining cases

B PULMONARY STENOSIS Three cases (19 to 21, Table III) Sinus rhythm signs suggesting right atrial enlargement, a QRS

axis almost +110 degrees, and lack of deep Q waves in Leads III and/or aV_F characterized all three cases (Fig 9) Essentially negative morphologies in Lead aV_R existed in 2 cases Initial QRS vectors abnormally directed towards the left anteriorly could be detected in all 3 cases Moreover in case 21, an abnormal q wave was present in Leads V_r and V_i RS complexes in the entire precordium (isodiphasism) were found only in case 19 (Fig 9) An rS pattern in the precordial leads was noted in case 20

Digitalis induced ST T changes in the left precordial leads were seen in one case In the remaining two the presence of these same changes could not be attributed to digitalis intake Right ventricular hypertrophy was diagnosed in case 21 with biventricular dilatation in case 19 (Fig 9)

III Group I, Subgroup C D loop

NO PULMONARY STENOSIS One case (22,

| Atrial enlargement | A QRS | Deep-Q III-aVr | aV _R | Isodiphasism | ST T changes | Ventricular pattern |
|--------------------|-------|----------------|-----------------|--------------|--------------|---------------------|
| RAH | +100 | + | QS | - | + | RVH |
| RAH | +80 | - | QS | + | + | BVH |
| RAH | +110 | + | Qr | + | - | RVH |
| RAH | +110 | - | QS | + | + | LVH |
| RAH | +10 | - | QS | + | + | BVH |
| RAH | 0 | + | Qr | - | - | LVH |
| RAH | +115 | + | QS | rS | + | LVH |
| RAH | +70° | - | QS | + | + | BVH |
| RAH | +120 | - | QS | + | - | BVH |
| RAH | +90 | + | QS | - | + | LVH |
| RAH | +110 | - | QS | + | + | BVH |
| RAH | +115 | - | Qr | rS | + | - |
| RAH | +110 | - | QR | - | + | RVH |

| Deep Q III-aVr | aV _R | Isodiphasism | ST T changes | Ventricular pattern | B V loop pulmonary outflow tract |
|----------------|-----------------|--------------|--------------|---------------------|----------------------------------|
| - | QR | + | + | BVH | D Loop—without PS |
| + | QS | + | - | - | L Loop—with P A * |

Sorted by a m acular co

Table IV) Briefly the most outstanding features were sinus rhythm right atrial enlargement undertermined QRS axis normal initial QRS forces isodiphasism digtals induced ST T changes and biven tricular hypertrophy Existing QR complex was seen in lead aV_R

IV Group I Subgroup C I loop

PULMONARY ATRESIA One case (23 Table IV) The most remarkable electrocardiographic findings of this case were a sinus rhythm right atrial hypertrophy a QRS axis of +90 degrees Deep Q waves in Leads III and aV_R a QS complex in Lead aV_R an initial QRS vector abnormally directed towards the left anteriorly and isodiphasism There were no signs of enlargement of any ventricle

V Group I Subgroup with Dextrocardia L loop

PULMONARY STENOSIS Three cases (24 to 26 Table V) Sinus rhythm in every case

Biatrrial enlargement in two cases right atrial hypertrophy being found in the remainder In case 25 the QRS axis was -90 degrees The QRS complex in Lead aV_R was essentially negative in two cases (24 and 26)

Right ventricular hypertrophy was diagnosed in one case Left ventricular enlargement was present in cases 24 and 26

The other electrocardiographic features summarized in Table V are not considered to be of any particular interest

VI Group II Subgroup A D loop

TWO CASES (27 and 28 Table VI) The most important electrocardiographic findings in these cases were the presence of an accentuated right ventricular enlargement with initial q waves in the right precordial leads (Fig 10)

VII Group II Subgroup B D loop

TWO CASES (29 and 30 Table VI) The QRS axis was of -90 degrees and ≈180

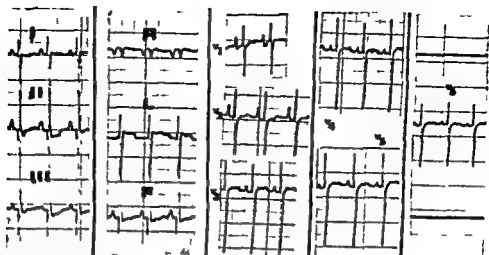


Fig 5 Electrocardiogram of Case 15. The main features are right atrial hypertrophy, deep Q waves in Lead III, an essentially negative deflection in aVR, ST-T changes, and rS pattern over the entire precordium (age = 3 months).

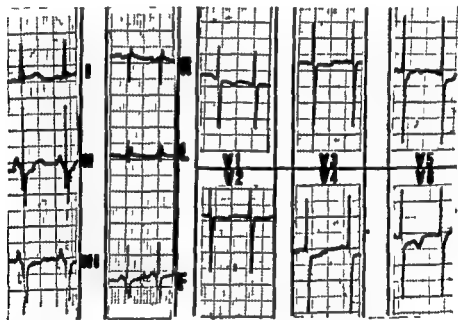


Fig 6 Deep Q waves, isodiphasism, and ST-T changes. Electrocardiogram of Case 14 (age = 4 months).

degrees in the two cases, respectively, both with a common atrioventricular orifice. Right atrial and ventricular enlargement were present in both cases.

VIII Group II, Subgroup C, D loop
ONE CASE (32, Table VI). The most important findings in this case were an undetermined QRS axis and biventricular hypertrophy. This latter correlated well with the ventricular proportions of this heart, the size of both, left and right ventricle being fairly similar.

Special case

This case, the rather unusual anatomic findings of which are displayed above, was

a heart in D Loop without pulmonary stenosis and two independent atrioventricular orifices. The most important electrocardiographic feature was the existence of biventricular hypertrophy with more conspicuous signs of left ventricular enlargement, a factor closely parallel to the anatomic ventricular proportions.

Discussion

In accordance with our expectations and as our findings confirmed, there are distinct electrocardiographic differences^{2,3} between the morphologies of Groups I and II; these differences will now be discussed. Such clear cut and extreme differences as are seen

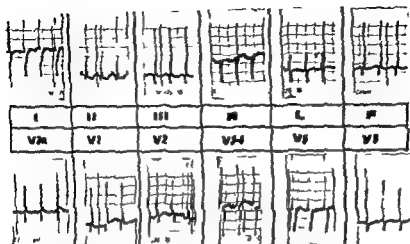


Fig 7 Electrocardiogram of Case 13. The most outstanding findings are the presence of deep Q waves in Lead V_1 and V_2 , ST-T changes and a QS pattern in Lead aV_R (age = 5 months)

in Figs 8 and 10 however are not the general rule in the cases concerned

Right arm morphologies have proved a reliable source of information in establishing the differential diagnosis between Groups I and II when the remaining electrocardiogram does not reveal the obvious differences shown in Figs 8 and 10. Thus essentially downward potentials in Lead aV_R are recorded in 86.9 per cent of our cases in Group I, particularly in those pertaining to Subgroups A and B (Tables II, III, IV and V). On the contrary, important upright deflections have been found in most of the cases of Group II (Table VI).

Upon having established and dealt with the major differences between Groups I and II cases, the individual findings pertaining to each of these will be treated separately.

Group I. In good agreement with the anatomic findings, a left ventricular enlargement existed in 66.6 per cent of the D loop cases, an isolated right ventricular hypertrophy being diagnosed only in L loop cases. Similar^{1,10} and different^{20,21} findings have been reported by others.

Biventricular hypertrophy (isodiphasic m) found in 34.7 per cent of the material was not related to any particular subgroup. The isodiphasism present in 47.8 per cent of our patients and more frequently in L loop cases has already been reported as characteristic in this group of anomalies.^{10,21}

In four L loop cases an unexpected left ventricular enlargement was present. It is

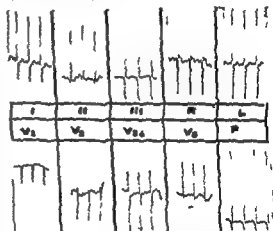


Fig 8 Case 18. Observe the pure left ventricular enlargement with a QS in Lead aV_R and ST-T changes in the left precordial leads (age = 4 months)

considered whether an almost antero-posterior relation of the infundibular chamber to the main ventricular cavity combined with unavoidable inexact position of the electrodes might be the origin of such paradoxical findings.

An rS pattern in the entire precordium was found in four patients. The incidence of this pattern was higher in cases with a D loop and normally related great arteries (as in two of our cases) and has been reported previously.¹¹ In one of the latter cases the height of the r wave increased after the creation of a systemic pulmonary artery anastomosis.

Table V Group I—Dextrocardia—ECG findings

| Case No and initials | Sub group | Age | Diagnosis | A V rates | G 4 * | Rhythm | Atrial enlargement |
|----------------------|-----------|------------------|-----------|-----------|-------|--------|--------------------|
| 24 J V D | A | 8 days | N | 2 | L TGA | S | RAH |
| 25 F G R | ? | 2 years | A | ? | L TGA | S | BAH |
| 26 P P V | ? | 8 months/4 years | A | ? | L TGA | S | BAH |

*See Table II for abbreviations.

Table VI Group II—ECG findings

| Case No and initials | Sub group | Age | Diagnosis | A V rates | G A | Rhythm | Atrial enlargement |
|----------------------|-----------|----------|-----------|-----------|--------|--------|--------------------|
| 27 S J | A | 4 months | N | CAVC | L TGA | S | LAH |
| 28 I E S | A | 19 days | N | 2 | D TGA | S | — |
| 29 N R | B | 15 days | N | CAVC | Normal | S | RAH |
| 30 P F B | B | 3 months | N | CAVC | D TGA | S | RAH |
| 31 I A | ? | 4 months | A | ? | D Malp | S | RAH |
| 32 R G D | C | 8 days | N | 2 | D TGA | S | — |
| 33 A G G | SC | 1 month | N | 2 | D Malp | S | BAH |

*Abbreviations LP = left posterior LA = left anterior RA = right anterior SC = special case For other abbreviations see Table II and

The essentially negative deflections in Lead aV_R observed in 86.9 per cent of our material and previously by others,¹⁹ seems to be the result of the right ventricular sinus hypoplasia.

A right atrial enlargement was present in 73.9 per cent of the cases. Batrial enlargement was present in 8.7 per cent all of which presented increased pulmonary blood flow, as previously mentioned.^{19, 21, 22, 25}

A left atrial hypertrophy has not been found in our material contrary to prior reports.^{23, 26, 27}

ST-T alterations in the left precordial leads were observed in 65.2 per cent of the cases. In 20 per cent these changes could not be attributed to digitalis intake. This factor, fairly common in our series, was found in cases with aortic obstruction.²⁸ We think that these alterations, whether brought about by digitalis intake or not, are the result of a volume overload of a left ventricular cavity receiving both the pulmonary and systemic venous flows.

Abnormal initial QRS forces directed toward the left anteriorly have been observed in 43.4 per cent of our material. As reported,^{19, 21, 22} these alterations were

more frequent in L loop cases. Nevertheless in our material the highest incidence was in cases associated with pulmonary stenosis (85.9 per cent) regardless of the nature of the bulbventricular loop.

The QRS was mostly located close to +120 degrees. In accordance with previous contributions,^{2, 18, 20} left axis deviation was more frequent in D loop cases.

As established,^{1, 22} the QRS axis and frontal and precordial morphologies were sometimes indistinguishable from those of tricuspid atresia cases.

In accordance with further series,^{2, 18, 20, 23, 29} other associated malformations did not significantly alter the QRS axis.

The earlier finding of discord between the QRS axis and the pattern of ventricular hypertrophy^{19, 20, 23, 29} did not prove statistically significant in our series as it was applicable only in three D loop and in two L loop cases.

Deep Q waves in Leads II, III, and aV_F were present in 30.4 per cent of the cases. Similar to prior recordings,^{23, 29} these were more frequent in L loop cases.

An atrioventricular block advanced

| QRS | Deep Q III aVF | aV _R | Isodisphatizm | ST T changes | Ventricular pattern | B V loop pulmonary outflow tract |
|---------|-------------------|-----------------|---------------|-----------------|------------------------|-------------------------------------|
| 1 +120 | — | QS | — | + | LVH | L Loop—with P A |
| 2 - 90 | — | qR | — | — | RVH | L Loop—with P S |
| 3 + 10° | — | rS | rS | — | LVH | L Loop—with P S |

| QRS | Deep Q III aVF | aV _R | Initial vector of QRS | Isodisphatizm | ST T changes | Ventricular pattern | B V loop pulmonary outflow tract |
|------|-------------------|-----------------|--------------------------|---------------|-----------------|------------------------|-------------------------------------|
| 80 | — | qR | LP | — | + | RVH | D Loop—without P.S |
| 160 | — | qR | LP | — | + | RVH | D Loop—with P S |
| 90 | — | qR | LA | + | + | RVH | D Loop—without P S |
| 180 | — | qR | LA | — | — | RVH | D Loop—with P S |
| +150 | — | qR | LP | — | — | RVH | D Loop—with P S |
| und | — | QR | LA | + | + | RVH | D Loop—without P S |
| +150 | — | QR | RA | — | + | RVH | D Loop—without P S |

degree (Fig 4) was observed in one L loop case. This has appeared more frequently in L loop cases 11 20 21 22.

Accelerated atrioventricular conduction was found in three cases two of which presented with a discordance between the position of the aorta (anterior and right sided respective of the pulmonary artery) and the type of bulboventricular loop (L loop).

Summing up we believe that single left ventricle cannot be electrocardiographically diagnosed. Nevertheless an electrocardiogram showing an essentially negative QRS morphology in Lead aV_R, ST T alterations in the leads facing the left ventricular free wall and stereotyped QRS morphology (RS or rS) in the entire precordium make this anomaly suspect. A typical patent ductus arteriosus, transposition of the great arteries with a large systemic shunt, tricuspid atresia and myocardial pathies must be considered in the differential diagnosis.

In cases with increased pulmonary blood flow P waves are generally taller in tricuspid atresia than in single left ventricle. There is a higher incidence of left atrial hypertrophy in myocardial pathies and in

patent ductus arteriosus in relation to single left ventricle.

Group I Dextrocardia. A QRS morphology inversion between Leads I and aV_R was found in the dextrocardiac group. In contrast a similar inversion has not been found in L loop cases with the cardiac apex pointing towards the left. The existence of P waves suggesting biatrial enlargement is worth emphasizing despite the presence of severe obstruction to the pulmonary outflow tract.

Group II. The electrocardiographic findings in this group were in good accord with the relative ventricular size: a right ventricular hypertrophy having been diagnosed in Subgroup A and B cases. There was a biventricular enlargement in Group C and in the special case. The QRS axis was uniformly directed towards the right.

In contrast to Group I an essentially positive deflection in Lead aV_R prevailed in this group.

The abnormally posterior and leftward orientation of the initial QRS forces in these cases (87.5 per cent) was attributed to an accentuated right ventricular hypertrophy.

We suggest that when considering a

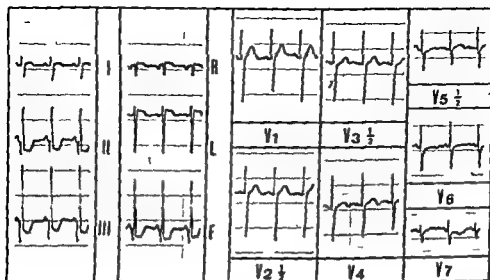


Fig 9 Electrocardiogram showing a QS complex in aVR, RS complexes through V₁ to V₆ and ST-T changes in Leads II, III, aVR, and the precordial leads. Case 19 (age = 1 day)

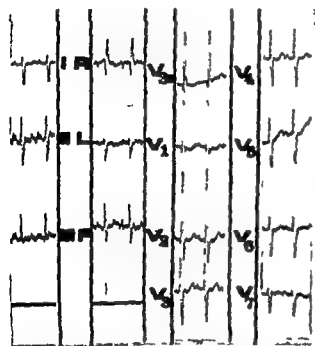


Fig 10 Accentuated right ventricular enlargement in one IIA case. Note the presence of q waves through V₁, V₂, and V₃ and the important terminal forces in Lead aVR. Case 28 (age = 19 days)

electrocardiogram showing an accentuated right ventricular enlargement with an essentially upward deflection in Lead aVR and q waves through V₁, V₂, and V₃; the diagnosis of the single right ventricle should be taken into account. Other conditions having a bearing on the differential diagnosis would include mitral atresia, double outlet right ventricle, severe right ventricular outflow obstructions, and total anomalous pulmonary venous drainage.

Summary

A review is made of the electrocardiographic findings of 33 cases with derangement in the spatial relationship between the ventricular septum and the atrioventricular canal.

To obtain an analysis with a maximal homogeneity six groups are established according to the extent and direction of the above mentioned derangement (Table I). Groups IA and IIA are equivalent to the most frequent designations of single left and right ventricle respectively or more exactly to those of double inlet left and right ventricle.

Cases with disturbance in the position of the heart are considered separately to ascertain whether the outflow obstructions and the bulboventricular loop influence the electrocardiographic morphologies in any way.

Electrocardiographic features of the other four groups (Table I) are analyzed to determine if the differences in the relationship between the atrioventricular canal and the interventricular septum are of small degree and therefore if the hypoplastic cavity becomes larger.

REFERENCES

1. Neill C A and Brink A J. Left axis deviation in tricuspid atresia and single ventricle: the electrocardiogram in 36 autopsied cases. *Circulation* 12:612, 1955.
2. Van Praagh H, Ongley J, and Swan

- II J C Anatomic types of single or common ventricle in man Morphologic and geometric aspects of 60 necropsied cases *Am J Cardiol* 23 367 1964
- 3 Van Praagh R Van Praagh S Vlad P and Keith J D Diagnosis of the anatomic types of single or common ventricle *Am J Cardiol* 15 315 1965
- 4 De la Cruz M V and Miller II L Double inlet left ventricle Two pathological specimens with comments on the embryology and on its relations to single ventricle *Circulation* 37 747 1968
- 5 Lev M Liberthson R R Kirkpatrick J R Friedrich A O Eckner O A F and Arcilla R A Single (primitive) ventricle *Circulation* 39 577 1969
- II Munoz Castellanos J Rodriguez Llorian A Martinez Rios M A and Espino Vela J Doble camara de entrada y salida del ventriculo derecho *Arch Inst Cardiol Mex* 39 114 1969
- 7 Liberthson R R Paul M H Muster A J Arcilla R A Eckner O A F and Lev M Straddling and displaced atrioventricular orifices and valves with primitive ventricles *Circulation* 43 213 1971
- 8 Quero Jimenez M Atresia of the left atrioventricular orifice associated with a Holmes heart *Circulation* 44 739 1970
- 9 Quero Jimenez M Coexistence of single ventricle with atresia of one atrioventricular orifice *Circulation* 46 794 1972
- 10 Lambert E C Single ventricle with a rudimentary outlet chamber Case report *Bull Johns Hopkins Hosp* 68 231 1951
- 11 Rastelli C G Ongley P A and Titus J L Ventricular septal defect of atrioventricular canal type with straddling right atrioventricular valve and mitral valve deformity *Circulation* 37 816 1968
- 12 Navarro-López F Marin J Zomeno M and Llorian A C Mitral atresia and occlusive left atrial thrombus A case with 11 years of survival *Br Heart J* 31 619 1969
- 13 Quero Jimenez M Trastornos en la relación del canal atrioventricular con el tabique inter ventricular Tesis doctoral Madrid 1971 (To be published)
- 14 Quero Jimenez M Perez Martinez V M Mañe Azcarate M J Merino Batres G and Moreno Granado F Exaggerated displacement of the atrioventricular canal toward the bulbus cordis (Rightward displacement of the mitral valve) *Br Heart J* 35 65 1973
- 15 Ziegler R F Electrocardiographic studies in normal infants and children Springfield Ill 1951 Charles C Thomas
- 16 Sod Pallares II Briten A and Medrano G A Electrocardiografía y vectocardiografía deductivas Mexico City 1964 La Prensa Médica Mexicana
- 17 Walsh S Z Electrocardiography in infants and children in Watson H ed *Pediatric cardiology* Saint Louis 1968 The C V Mosby Company
- 18 Van Praagh R Single (common) ventricle in Keith J D Rowe R D and Vlad P editors *Heart disease in infancy and childhood* New York 1967 Macmillan Publishing Co Inc
- 19 Burch G E and DePasquale N P Electrocardiography in the diagnosis of congenital heart disease Philadelphia 1967 Lea & Febiger Publishers
- 20 Shaher R M The electrocardiogram in single ventricle *Br Heart J* 25 465 1963
- 21 Davachi F and Molier J A The electrocardiogram and vectocardiogram in single ventricle Anatomic correlations *Am J Cardiol* 23 19 1969
- 22 Nadas A S *Pediatric cardiology* Philadelphia 1957 W B Saunders Company
- 23 Elliott L P Ruttenberg H B Elliot R S and Anderson R C Vectorial analysis of the electrocardiogram in common ventricle *Br Heart J* 26 302 1964
- 24 Keith J D Rowe R D and Vlad P *Heart disease in infancy and childhood* New York 1958 Macmillan Publishing Co p 512
- 25 Anselmi C Armas S M De la Cruz M V De Pisani F and Blanco P Diagnosis and classification of single ventricle *Am J Cardiol* 21 813 1968
- 26 Gasul M B Arcilla R A and Lev M II S *Heart disease in children* London and Philadelphia 1966 Pitman Medical Publishing Co Ltd J B Lippincott Co
- 27 Honey M Rushton D G and Taylor D G Single ventricle with rudimentary chamber *Guys Hosp Rep* 109 116 1960
- 28 Engle M A in Watson S *Pediatric cardiology* Saint Louis 1968 The C V Mosby Company p 632
- 29 Taussig H B *Congenital malformations of the heart* 2nd ed vol II Cambridge Mass 1960 Harvard University Press
- 30 Morgan A D Kroyetz L J and Shiebler G L Electrocardiographic analysis of nine cases of single ventricle with the great vessel arrangement of congenitally corrected transposition in *Vectocardiography* 1965 Amsterdam 1966 North Holland Publishing Company p 327
- 31 Elliott L P and Morgan A II in Moss A J and Adams F H *Heart disease in infants children and adolescents* Baltimore 1968 The Williams & Wilkins Company p 589

Radiological patterns of obstructive cardiomyopathy of the left ventricle in childhood

C Pernot, MD

J C Hocffel MD

M Henry, MD

A M Worms, MD

G Rothhahn, MD

Dommartin Les Toul France

Sub-aortic muscular stenosis or obstructive cardiomyopathy of the left ventricle is a well known disease in adults but few reports concern children and infants. We have collected nine cases four of which were in infants of 3, 6, 11 and 18 months, and five in children of 3, 4, 5, 5½ and 6 years respectively.

Obstructive cardiomyopathy is generally idiopathic and often familial. It is sometimes associated with stenosis of the aortic outflow tract. A systolic murmur was present in all cases. In infants cardiac failure may be the first symptom. In children the murmur may be discovered at routine examination or the child may present with dyspnea and even syncope in some cases. The systolic murmur is ejection like, variable and best heard at the apex. The ECG may show signs of left ventricular hypertrophy or of biventricular hypertrophy if the cardiopathy is bilateral. We have observed 6 cases of primary cardiomyopathy of which 2 in infants were biventricular, and 3 cases of secondary cardiomyopathy in children, of the latter one was in a case of aortic valvular stenosis.

the second a case of subvalvular (diaphragmatic) aortic stenosis, and the third a case of both valvular and subvalvular stenosis.

The frontal view (Table I)

In the 4 cases in infants the cardiothoracic index was above 0.55, the mean at this age. In the 5 cases in children the index is superior to the normal mean of 0.45. In all cases there was cardiomegaly. The inferior left cardiac segment was convex and the apex was displaced laterally (Fig. 1). These signs of left ventricular hypertrophy were noted in 8 out of 9 cases. Angulation of the left superior border as described by Wigle and Vernant was seen in only one case, an infant of 6 months.

Angiocardiography

A ventricular systolic pressure gradient was present in all cases. Angiocardiography and left cineangiography were performed with a catheter introduced by way of a retrograde arterial route. Frontal, right anterior oblique and lateral views were taken. The left ventricle looks abnormal being distorted with a very thick wall.

From the Services de Cardiologie et de Radiologie de l'Hôpital Jeanne d'Arc

Received for publication Nov. 27, 1972

Reprint requests to Dr. J. C. Hocffel, Services de Cardiologie et de Radiologie de l'Hôpital Jeanne d'Arc 54 Dommartin Les Toul France

Table I Chest film—frontal view

| Patient no | Age | Cardiothoracic index | Apex | Left inferior segment of the heart |
|------------|-----------|----------------------|--------------------------|------------------------------------|
| 1 | 18 months | 0.62 | much displaced laterally | convex |
| 2 | 3 months | 0.62 | much displaced laterally | markedly convex |
| 3 | 6 months | 0.63 | much displaced laterally | markedly convex |
| 4 | 11 months | 0.64 | displaced laterally | markedly convex |
| 5 | 6 years | 0.60 | displaced laterally | normal |
| 6 | 3 years | 0.60 | displaced laterally | convex |
| 7 | 3½ years | 0.52 | displaced laterally | markedly convex |
| 8 | 5 years | 0.48 | not displaced | convex |
| 9 | 4 years | 0.54 | displaced laterally | markedly convex |



Fig 1 Typical chest x ray of infant with cardiomyopathy. The inferior left cardiac segment is convex and the apex is laterally displaced.



Fig 2 Border abnormalities in 11 month old infant with abnormal left ventricle. There is an indentation of the left border of the upper portion of the left ventricle and also of the inferior margin associated with septal hypertrophy.

A The morphologic or qualitative aspects

1 THE ABNORMALITIES OF THE BORDERS

We noted an indentation of the left border of the upper part of the left ventricle and of the inferior margin in 7 out of the 9 cases (Fig 2). These indentations were associated with septal hypertrophy and were visible in both diastole and systole. The apex was not delineated or was to a lesser degree tapered in 5 out of the 9 cases.

2 THE INTERNAL APPEARANCE OF THE VENTRICULAR CAVITY

Hypertrophy of the mitral papillary muscles is usual (Figs 2 and 3). The anterior one looked very hypertrophied in 6 cases and the posterior muscle looked very hypertrophied in 7 cases out of the 9.

This sign is not specific but is very common and is easily visualized. If the axis of the papillary muscles is displaced to the

left or upward instead of being continuous with the mitral axis, this is of high diagnostic value. This distortion of the papillary muscles causes abnormal traction on the chordae tendineae and the mitral valve anterior leaflet cannot move backward during systole. The outflow tract is obstructed during meso- and telesystole.

The mid ventricular stenosis in systole is evident in 7 cases out of 9 cases (Fig 3). The ventricular cavity looks very small during telesystole in these cases and is

Radiological patterns of obstructive cardiomyopathy of the left ventricle in childhood

C Pernot, MD

J C Hoeffel, MD

M Henry, MD

A M Worms, MD

G Rothhahn MD

Dommartin Les Toul, France

Sub-aortic muscular stenosis or obstructive cardiomyopathy of the left ventricle is a well known disease in adults but few reports concern children and infants. We have collected nine cases four of which were in infants of 3, 6, 11, and 18 months, and five in children of 3, 4, 5, 5½, and 6 years respectively.

Obstructive cardiomyopathy is generally idiopathic and often familial. It is sometimes associated with stenosis of the aortic outflow tract. A systolic murmur was present in all cases. In infants cardiac failure may be the first symptom. In children the murmur may be discovered at routine examination, or the child may present with dyspnea and even syncope in some cases. The systolic murmur is ejection like variable, and best heard at the apex. The ECG may show signs of left ventricular hypertrophy or of biventricular hypertrophy if the cardiopathy is bilateral. We have observed 6 cases of primary cardiomyopathy of which 2 in infants were biventricular, and 3 cases of secondary cardiomyopathy in children of the latter, one was in a case of aortic valvular stenosis

the second a case of subvalvular (diaphragmatic) aortic stenosis and the third a case of both valvular and subvalvular stenosis.

The frontal view (Table I)

In the 4 cases in infants the cardiothoracic index was above 0.55, the mean at this age. In the 5 cases in children the index is superior to the normal mean of 0.45. In all cases there was cardiomegaly. The inferior left cardiac segment was convex and the apex was displaced laterally (Fig 1). These signs of left ventricular hypertrophy were noted in 8 out of 9 cases. Angulation of the left superior border as described by Wigle and Vernant was seen in only one case, an infant of 6 months.

Angiocardiography

A ventricular systolic pressure gradient was present in all cases. Angiocardiography and left cineangiography were performed with a catheter introduced by way of a retrograde arterial route. Frontal, right anterior oblique and lateral views were taken. The left ventricle looks abnormal being distorted with a very thick wall.

From the Services de Cardiologie et de Radiologie de l'Hôpital Jeanne d'Arc.

Received for publication Nov. 27, 1972.

Reprint requests to Dr J. C. Hoeffel, Services de Cardiologie et de Radiologie de l'Hôpital Jeanne d'Arc 54 Dommartin Les Toul, France.

October 1973 Vol. 4, No. 4, pp. 462-466

Table II Thickness of the left ventricular wall

| Manifestation | Diastolic thickness (mm) | Systolic thickness (mm) | Systolic diastolic difference (mm) | Diastolic thickness | Systolic thickness | Cavity width | Cavity width |
|-----------------------|--------------------------|-------------------------|------------------------------------|----------------------|----------------------|--------------------------|-------------------------|
| | | | | Aortic diameter (mm) | Aortic diameter (mm) | Diastolic thickness (mm) | Systolic thickness (mm) |
| cardiomyopathy | 8.6 | 17.6 | 7.8 | 5.62 | 11.5 | 0.51 | 0.17 |
| normal left ventricle | 5.5 | 8.5 | 3 | 3.19 | 4.95 | 0.94 | 0.52 |



Fig 5 Angiocardiogram showing mitral regurgitation in an 18-month-old child. This symptom was present in two of our patients.

infants 10.2 mm in the children 3 to 6 years of age) compared with 5.5 mm for normal ventricles. During systole the thickness of the left ventricle was 16.6 mm in the cardiomyopathies (15 mm in the infants and 19 mm in the children 3 to 6 years of age) compared with 8 mm for normal ventricles. In order to remove the factor of age we calculated the relationship between the mean diastolic thickness and the aortic diameter. The same relationship was calculated for the systolic thickness. In diastole the mean ratio was 5.62 (normal 3.19) and in systole 11.5 (normal 4.95). Finally the relationship between the difference between mean diastolic and mean systolic thickness and the aortic diameter was 5.9 in the cardiomyopathies compared with 1.74 for normal ventricles. In their series of adults Levine and Braunwald



Fig 6 Angiocardiogram of 18-month-old infant with cardiomyopathy. Infundibular stenosis is apparent on the left with taper in the dilated inflow tract.

studied the relationship between the width of the ventricular cavity and the wall thickness and noted that it was much smaller in cardiomyopathies than in valvular stenosis. Our own results show values of 0.51 in diastole and 0.17 in systole for cardiomyopathies whereas measurements on normal ventricles gave values of 0.94 and 0.52 respectively.

In conclusion angiography of the left ventricle in infants and children with cardiomyopathy show similar signs to those found in adults. The left ventricle is distorted with indentation of the margins and



Fig 3 Angiocardiogram showing the internal appearance of the ventricular cavity. Hypertrophy of the mitral papillary muscles is evident and mid ventricular stenosis in systole is also evident.



Fig 4 Angiocardiogram lateral view of a 6-year old child showing the pattern of an inverted cone which appeared very distinctly in two of our patients. This manifestation is very diagnostic if seen in telesystole.

moderately diminished in size during diastole.

The characteristic feature of a radiolucent band crossing the outflow tract below the aortic ring, visible on frontal and lateral views in mesosystole and in telesystole was difficult to see in most of our cases.

On lateral views we noted the pattern of an inverted cone (Fig 4) which was very distinct in two out of six cases clear in one, and absent in three cases. If seen in telesystole it is highly diagnostic. The cone is limited anteriorly by the hypertrophied septum and posteriorly by the anterior mitral leaflet which does not move backward during systole. It can also be seen in diastole but must be differentiated from the false image of stenosis in diastole secondary to the opening of the anterior leaflet which comes close to the septum. Finally, we noted mitral regurgitation in 2 cases (Fig 5).

B The quantitative aspects

We have tried to measure some of the

abnormalities, in particular the thickness of the left ventricular wall. Firstly the longitudinal axis of the ventricle is altered and an angulation of 167 degrees (mean value) was seen as compared with 119 degrees for 9 normal ventricles (in infants and children of the same age). The reduction of the systolic capacity is marked at the middle segment of the left ventricle in both absolute and relative terms. In order to establish this fact we have measured the relationship comparing the difference between the diastolic and systolic widths of the ventricular cavity with the diameter of the aorta. In normal ventricles this ratio was 0.41 but in the patients with obstructive cardiomyopathy the mean value was 0.81. The thickness of the left ventricular wall at the middle section of the free left margin on frontal views appeared much greater in the cardiomyopathies than in the normal left ventricles (Table II). During diastole the mean thickness was 8.6 mm in the cardiomyopathies (7 mm in the

Embolic coronary artery occlusion in percutaneous transfemoral coronary arteriography

Angel de la Torre MD*

Daniel Jacobs MD**

Juan Aleman MD***

George A Anderson MD****

Jacksonville Fla

Although the technique of selective coronary angiography as described by Sones and Shirey¹ has been in popular use since 1962 and has enjoyed a record of relative freedom from major complications its few limitations including occasional poor coronary opacification difficulty with catheter introduction associated with anatomic arterial variations or the inherent complexity of the procedure together with the advent of coronary bypass surgery with the resultant demand for increasing numbers of studies have led to a search for newer more reliable techniques

Among the alternatives the technique of percutaneous transfemoral artery bilateral selective coronary arteriography as described by Judkins² has indeed seemed to resolve most of the problems previously encountered

With a recent exception³ previous reports of a major complication of coronary artery

occlusion with myocardial infarction due to emboli introduced with this technique did not prepare us for its startling frequency in our laboratory^{2,4,5}

Methods and materials

Selective coronary arteriography was carried out in 215 patients from April 1970 to May 1972 (147 men and 68 women whose ages ranged from 30 to 70 years with a mean age of 52 years) The Sones technique was used in 76 and the Judkins technique was used in 139 During the Judkins technique a No 18 gauge thin walled arterial needle was employed for percutaneous entry of the common femoral artery Teflon coated 145 cm guide wires were used and where necessary a Safety J guide was employed

A short 7 F Teflon catheter was inserted into the artery over the guide before inserting the pig tail catheter used for the

From the Department of Medicine University Hospital Jacksonville, Fla.

Supported in part by a grant from the Northeast Florida Heart Association.

Received for publication December 4, 1972

Reprints: Dr. George A. Anderson, 2005 E. 7th Ave., Jacksonville, Fla. 32204

Associate Professor of Medicine, Division of Cardiology JHEP University of Florida, University Hospital, Jacksonville, Fla. Chief, Coronary Catheterization Laboratory St. Vincent's Hospital Jacksonville, Fla.

Assistant Professor of Medicine, Division of Cardiology JHEP University of Florida, University Hospital Jacksonville, Fla.

Professor of Cardiology, University Hospital Jacksonville, Fla.

Clinical Associate Professor of Medicine, Division of Cardiology JHEP University of Florida, University Hospital, Jacksonville, Fla.

a reduced systolic capacity. The thickness of the free wall of the left ventricle is increased and in infants may exceed 5.5 mm in diastole and 8.5 mm in systole. In the biventricular cases, right ventricular anomalies may also be seen. The right ventricle is usually displaced because of the septal hypertrophy and enlargement of the left ventricle. In infants, infundibular stenosis may be seen on the right with stasis in the inflow tract which is often dilated (Fig 6). In these biventricular cases the thickness of the septum may be measured by means of simultaneous opacification of both ventricles with a right catheter and 2 separate injections (see Robicsek and associates).

Differential diagnosis

In children cardiomyopathy may be confused with muscular hypertrophy of the left ventricle as seen in Friedrich's ataxia or with the tumor like hypertrophies associated with tuberous sclerosis or Bourneville's glycogen storage disease or Refsum's disease. The clinical picture may be more valuable than radiology in differentiating the hypertrophic cardiomyopathies from the obstructive cardiomyopathies either idiopathic or secondary to a congenital stenosis of the aortic outflow tract.

Summary

The authors report 9 cases of obstructive cardiomyopathy of the left ventricle in childhood. Angiograms performed by the Seldinger technique showed the characteristic radiological features of the disease which may be either idiopathic in origin or associated with an aortic stenosis.

REFERENCES

- Bourdarias J P, Penher Ph, Ourbak P and Lenegre J L. Myocardio-pathie obstructive. *Rev Prat* 17:228 1967.
- Braunwald E, Lambrew C T, Rockoff S D, Ross J Jr and Morrow A G. Idiopathic hypertrophic subaortic stenosis: a description of disease based upon an analysis of 64 patients. *Circulation* 29 and 30 (Suppl IV) 3 1964.
- Cohen J, Effat H, Goodwin J F, Dakley C M and Steiner R E. Hypertrophic obstructive cardiomyopathy. *Br Heart J* 26:16 1964.
- Dodge H T. Determination of left ventricular volume and mass. *Radiol Clin North Am* 18:459 1971.
- Ferrane J, Bourdarias J P, Lockhart A, Ourbak P, Seebat L and Lenegre J. Cinéangiographie dans les myocardio-pathies obstructives. *Arch Mal Coeur* 5: 739 1964.
- Gerbaux A, Penher Ph and Gillet J L. Les sténoses musculaires des ventricules. *Rev Prat* 20:241 1970.
- Geschwind H. Apport de l'angio et de la cinéangiographie dans le diagnostic des obstacles à l'éjection ventriculaire. *Gaz Méd de France* 77:2909 1970.
- Greene D G, Carlisle R, Grant C and Bunnell I L. Estimation of left ventricular volume by one-plane cineangiography. *Circulation* 36:61 1967.
- Houplon M. A propos des cardiomyopathies obstructives. Medical Thesis Nancy 1967 283 pp.
- Levine N D, Rockoff S D and Braunwald E. An angiographic analysis of the thickness of the left ventricular wall and cavity in aortic stenosis and other valvular lesions. Hemodynamic angiographic correlations in patients with obstruction to left ventricular outflow. *Circulation* 28:339 1963.
- Moes C A F, Peckham G B and Keith J D. Idiopathic hypertrophy of the interventricular septum causing muscular subaortic stenosis in children. *Radiology* 83:283 1964.
- Moss A J and Adams F H. Heart diseases in infants, children and adolescents. Baltimore, 1968. The Williams & Wilkins Company, p 568.
- Robicsek F, Gatling H B, Daugherty H K and Mullen D C. Two x-ray signs helpful in the diagnosis of hypertrophic cardiomyopathy. *Am Heart J* 80:606 1970.
- Rothblat G. Etude radiologique du ventricule gauche dans les cardiomyopathies obstructives de l'enfant. Mémoire C E S. *Radiol Méd Nancy* 1972 96 pp.
- Rudhe U, Zetterquist P and Wallgren G. Angiocardiography of hypertrophic obstructive cardiomyopathy in infancy and childhood. *Ann Radiol* 9:185 1966.
- Shemtov A, Deutsch V, Yahini J H and Neufeld H N. Cardiomyopathy associated with congenital heart disease. *Br Heart J* 33:182 1971.
- Simon A L, Ross J Jr and Gault J H. Angiocardiographic anatomy of the left ventricle and mitral valve in idiopathic hypertrophic subvalvular stenosis. *Circulation* 36:857 1967.
- Soulié P, Forman J, Delzant J F, Dupenier C and Vain J. Sténose musculaire du ventricule gauche. Hémodynamique et phonocardiographie intracardiacque (a propos de 35 cas). *Arch Mal Coeur* 60:1 1967.
- Vernant P and Sterba S. Cardiomyopathie obstructive. Signes cliniques-physiopathologiques-traitement. *Gaz Méd de France* 77:2933 1970.
- Watson H. Pediatric cardiology. London 1968. Lloyd Luke Ltd p 342.

71 m 86
mb r 4

and/or ischemic papillary muscle dysfunction she was admitted to the hospital in October 1971 for cardiac catheterization and selective coronary arteriography.

During selective injection of the left coronary artery using the percutaneous transfemoral approach she suddenly developed chest pain accompanied by cardiovascular collapse. She was found to have a block of the left main coronary artery at its bifurcation into the circumflex and anterior descending branches (Fig 1). Resuscitation attempts precluded recording her electrocardiogram at this particular time and despite all efforts the patient died on the catheterization table.

Postmortem examination revealed an atherosclerotic narrowing of greater than 80 per cent prior to the bifurcation of the left main coronary artery into the left anterior descending and circumflex branches which was occluded by a fresh clot. Findings compatible with the syndrome of ballooning mitral valve were present.

Patient 2 A 56-year old white man was first admitted to the Duval Medical Center in August 1969 presenting a six month history of progressively more severe and frequent substernal chest pain which radiated to the left arm and was usually relieved by rest.

Physical examination was within normal limits and after serial electrocardiograms and enzyme determinations were unremarkable he was discharged with a diagnosis of coronary insufficiency.

After readmission in April 1970 using the Sones technique bilateral selective coronary arteriography demonstrated 10 per cent narrowing of the right coronary artery and complete occlusion of the left anterior descending artery beyond the first diagonal branch with retrograde filling from the right coronary artery. In June 1970 an aorto-left anterior descending artery saphenous vein bypass procedure was performed successfully.

After one year free of anginal symptoms he was readmitted in August 1971 for restudy of his bypass graft. Using the percutaneous transfemoral approach injection into the bypass revealed graft patency but injection into his left main coronary artery was followed immediately by bradycardia and complete IV block accompanying embolic occlusion of the left main coronary artery (Figs 2 and 3). A pacemaker was positioned in the catheterization laboratory however after transfer he died in the Coronary Care Unit in cardiogenic shock. Permission for autopsy was not granted.

Patient 3 In January 1970 a 32 year old white man began experiencing severe substernal chest pain which radiated to both shoulders was occasionally accompanied by pallor, bore no relation to exertion and was rarely relieved by nitroglycerin. He smoked one package of cigarettes daily for 20 years and family history contained evidence of coronary artery disease.

In April 1970 after a normal admission physical examination percutaneous transfemoral selective coronary arteriography was carried out. On injection into the left coronary artery the distal left anterior descending artery seemed to be totally blocked about 2 cm beyond its origin (Fig 4). Accompanying the monitor revealed pronounced anterior



Fig 2 Patient No 2 Percutaneous transfemoral injection of the bypass graft left anterior oblique projection. Prior to injection of the left main coronary artery all three branches of the left coronary system were patent.



Fig 3 Patient No 2 Percutaneous transfemoral left coronary arteriography left anterior oblique projection. Injection into main left coronary artery demonstrated complete block of the left main coronary artery. (Catheter was retracted after injection.)

ST segment and T wave elevation and because of the persistent pain the procedure was terminated without injection of the right coronary artery.

He was transferred to the Coronary Care Unit and had an uneventful recovery from an anterior myocardial infarction confirmed by electrocardiogram but he continued to complain of substernal chest pain identical to that for which he was first admitted.

Patient 4 A 53 year old Negro male construction worker was admitted to the University Hospital in May 1972 presenting a history of precordial chest pain relieved within 10 minutes by nitroglycerin for approximately five years more frequent since December 1971. He had smoked more than two packages of cigarettes daily for more than 20 years before stopping 6 to 8 years prior to admission and was known to be hypertensive for 4 years treated sporadically. Physical examination disclosed a blood



Fig 1 Patient No 1 Percutaneous trans femoral left coronary arteriography left anterior oblique position. Injection of the left main coronary artery revealed total obstruction proximal to bifurcation into the left anterior descending and circumflex branches.

left ventriculogram. Dacor Judkins pre shaped disposable right and left coronary catheters were used for the coronary study.² Selective studies were done with the patient in different oblique projections. Contrast material was injected by hand syringe and pressures were monitored by the stop cock system described by Sones and Shurey.³ Flushing of the coronary catheter which was continuously infused with heparinized saline was made at the level of the descending aorta before it was advanced toward the ascending aorta. At the present time the left coronary catheter is removed from the coronary ostium after each injection aspirated, and is flushed before reinsertion. If right side cardiac catheterization is not performed an intravenous catheter is placed in the left arm before the procedure for use in emergency situations. Pressures were monitored with P23Db Statham pressure transducers and mean artery pressure was obtained by electronic filtering. Lead II of the electrocardiogram was continuously inscribed on all records. All tracings were inscribed by a 16 channel Electronics for Medicine DR 16 physiological Recorder.

Following the procedure the patients were admitted to the Coronary Care Unit for 24 hours during which time they were monitored continuously and vital signs were taken periodically. An electrocardio-

gram was taken the following day before the patient was discharged.

If, because of clinical angiographic electrocardiographic findings a cardiac complication was suspected serial enzymes and electrocardiograms were obtained.

Results

We have done 139 selective coronary studies via the percutaneous trans-femoral artery approach from April 1970, to May 1972. Approximately 1112 injections were made into the coronary arteries during these studies.

During the procedure acute myocardial infarction probably due to embolic occlusion within the left coronary artery system occurred in 6 patients. Of these patients No 1 and 2 died with total occlusion of the left main coronary artery shortly after the procedure.

Postmortem examination obtained in Patient 1 disclosed a fresh clot obstructing the left main coronary artery at the site of severe atherosclerotic narrowing. In the case of Patient 2 consent for autopsy was not obtained.

Patient 3 developed total occlusion 1 cm distal to the origin of the left anterior descending coronary artery after the first injection. Patient 4 also developed total occlusion of the left anterior descending coronary artery at the junction between its middle and distal thirds. Both patients experienced uneventful recovery from clinical evidence of acute myocardial infarction.

Patients 5 and 6 both suffered total occlusion of the left circumflex coronary artery. In both there was also uneventful recovery from clinical evidence of acute myocardial infarction.

Case reports

Patient 1 A 62 year old white woman diagnosed in 1967 as having Turner's syndrome (XO45 XX46 mosaic by chromosome analysis) was followed at the Duxon Medical Center since 1965 for mild congestive heart failure associated with chest pain relieved within 5 to 10 minutes by nitroglycerin.

In 1967 a short late systolic murmur with mid systolic click was described. Although the electrocardiogram was within normal limits, Masters test demonstrated subendocardial injury during, and immediately after exercise.

As she was felt to have ballooning mitral valve



Fig 7 Patient No 5 Percutaneous transfemoral left coronary arteriography right anterior oblique projection First injection demonstrated patency of all three branches of the left coronary system Note minimal disease of left anterior descending coronary artery



Fig 9 Patient No 6 Percutaneous transfemoral left coronary arteriography left anterior oblique projection First injection demonstrated all three branches of the left coronary artery system to be patent



Fig 8 Patient No 5 Percutaneous transfemoral left coronary arteriography right anterior oblique projection Fourth injection demonstrated block of left circumflex coronary artery



Fig 10 Patient No 6 Percutaneous transfemoral left coronary arteriography left anterior oblique projection Third injection demonstrated block of left circumflex coronary artery

In January 1972 following a normal physical examination percutaneous transfemoral bilateral selective coronary arteriography was carried out in which a mild right coronary artery was found occluded and the left anterior descending coronary artery was found to be mildly diseased. Following the third injection into the left main coronary artery he developed hypotension and bradycardia confirmed by electrocardiographic evidence of inferior wall injury. There was noted to be complete obstruction of the left circumflex coronary artery which had not been seen in the previous injections in the same projection (Figs 9 and 10). He was transferred to the Coronary Care Unit where his serum enzyme rose and electrocardiographic evidence of anterolateral myocardial infarction developed.

Discussion

We had used almost exclusively the technique of selective coronary arteriography described by Sones and Shirey¹ but in 1970 we started using the percutaneous transfemoral technique^{2,4} in the following situations

- 1 tortuous subclavian innominate arteries
- 2 difficulty in obtaining satisfactory coronary opacification
- 3 previous cut-downs in the brachial artery with resultant



Fig 4 Patient No 3 Percutaneous trans-femoral left coronary arteriography left anterior oblique projection. Injection into the left main coronary artery disclosed total block of left anterior descending coronary artery. It should be noted the left main coronary artery and the left circumflex coronary arteries are patent.

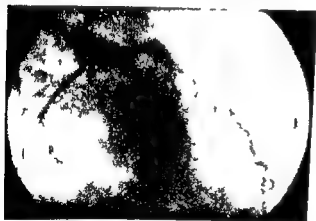


Fig 5 Patient No 4 Percutaneous trans-femoral left coronary arteriography right anterior oblique projection. Normal left coronary system.

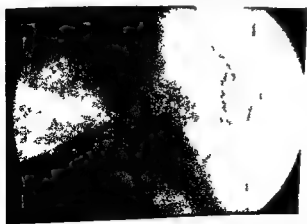


Fig 6 Patient No 4 Percutaneous trans-femoral left coronary arteriography right anterior oblique projection. Third injection demonstrated complete block of left anterior descending coronary artery.

pressure of 160/90 in accentuated though physiologically split second sound an audible fourth sound and a nonradiating Grade I aortic systolic murmur at the cardiac apex.

Having during bilateral selective coronary arteriography via the percutaneous transfemoral approach demonstrated normal right left main circumflex coronary arteries the patient developed substernal chest pain after the third injection into the left main coronary artery. Acute occlusion of the distal left anterior descending coronary artery was found (Figs 5 and 6) accompanied by ST segment elevation in the anterior and inferior leads followed later by evolving electrocardiographic changes.

Following the procedure he was admitted to the Coronary Care Unit and recovered uneventfully from an acute myocardial infarction due to embolic occlusion of the anterior descending branch of the left coronary artery.

Patient 5 A 56-year old white man was admitted to University Hospital in January 1972 for selective coronary arteriography having in 1960 begun experiencing substernal chest pain which radiated into the left arm occurred with exertion and anxiety and was relieved within 5 to 10 minutes by nitroglycerin.

For two and one half years he had had symptoms compatible with paroxysmal nocturnal dyspnea two or three pillow orthopnea and occasional exertional dyspnea during which time he was maintained on digitalis and diuretics. Hypertension known present for four years had been treated for three years. He had smoked 2 to 3 packages of cigarettes daily for 12 years stopping 10 years prior to admission. He was said to have had an elevated cholesterol and an abnormal glucose tolerance test. A brother and sister both 58 reportedly died of myocardial infarction and a brother 60 was said to have angina.

On physical examination blood pressure was 140/80 the optic fundi presented Grade II arteriosclerotic changes and there was an apical fourth sound.

On undergoing percutaneous transfemoral bilateral selective coronary arteriography in which the right coronary artery was within normal limits and the left anterior descending coronary artery displayed mild disease he sustained an embolus to the left circumflex artery on the fourth injection into the left main coronary artery (Figs 7 and 8). He was admitted to the Coronary Care Unit and after 14 hospital days was transferred to the Veterans Administration Hospital for continued convalescence from myocardial infarction.

The acute episode was unaccompanied by diagnostic electrocardiographic changes but infarction was reflected by marked rise in the serum enzyme.

Patient 6 A 52 year old white man was admitted to another hospital in December 1971 because of acute onset of substernal chest pain. Serial electrocardiograms demonstrated stable nonspecific ST T changes and after the usual studies ruled out acute myocardial infarction he was felt to have angina pectoris. Following discharge recurrent substernal pain brought on by exertion was partially relieved within 5 to 10 minutes by nitroglycerin. After institution of isosorbide and propranolol he became essentially symptom free.



Fig 7 Patient No 5 Percutaneous transfemoral left coronary arteriography right anterior oblique projection First injection demonstrated patency of all three vessels of the left coronary system Note minimal disease of left anterior descending coronary artery



Fig 9 Patient No 6 Percutaneous transfemoral left coronary arteriography left anterior oblique projection First injection demonstrated all three branches of the left coronary artery system to be patent



Fig 8 Patient No 5 Percutaneous transfemoral left coronary arteriography right anterior oblique projection Fourth injection demonstrated block of left circumflex coronary artery



Fig 10 Patient No 3 Percutaneous transfemoral left coronary arteriography left anterior oblique projection Third injection demonstrated block of left circumflex coronary artery

In January 1972 following a normal physical examination percutaneous transfemoral bilateral selective coronary arteriography was carried out in which a small right coronary artery was found occluded and the left anterior descending coronary artery was found to be mildly diseased. Following the third injection into the left main coronary artery he developed hypotension and bradycardia accompanied by electrocardiographic evidence of inferior wall infarction. There was noted to be complete obstruction of the left circumflex coronary artery, which had not been seen in the previous injections in the same projection (Figs 9 and 10). He was transferred to the Coronary Care Unit where his serial enzyme assays and electrocardiographic evidence of posterior wall small myocardial infarction developed.

Discussion

We had used almost exclusively the technique of selective coronary arteriography described by Sones and Shirey¹ but in 1970 we started using the percutaneous transfemoral technique^{2,4} in the following situations

- 1 tortuous subclavian innominate arteries
- 2 difficulty in obtaining satisfactory coronary opacification
- 3 previous cut-downs in the brachial artery with faint pulse

4 young females with spastic brachial arteries

However because the procedure is quickly learned by trainees fluoroscopic time is abbreviated, and good coronary visualization is obtained, its use in our laboratory had recently been greatly expanded

With one exception² our incidence of the procedural complication of myocardial infarction due apparently to embolic occlusion within the left coronary artery system appears to differ significantly from that reported by others though the problem may have been associated with left anterior descending artery occlusion described by Gau and colleagues³ and left circumflex coronary artery occlusion reported by Cheng and associates⁴

This potentially hazardous complication has, however been previously reported^{5,6,7} In order to minimize the risk it was first recommended that the catheter be energetically flushed prior to advancing the tip from the descending to the ascending aorta and, later if more than the usual time between the injections is required if persistent electrocardiographic changes occur or if the patient's position must be changed the catheter should be removed from the ostium of the coronary artery aspirated and flushed before another injection is carried out⁷

However, since two more cases have recently occurred in our laboratory despite meticulous observation of these modifications in technique we have elected to repeat this maneuver after each injection

The severity of the infarction apparently bears close relationship to the size of the embolus the level of the obstruction and the previous state of the coronary arteries. A small fibrin clot will probably not obstruct the main coronary artery or the initial segments of the primary branches unless there is a severe narrowing at these levels as was the case in Patients 1 and 2

In none of our patients was damping of the pressure noted prior to injection of contrast media into the left coronary artery—indicating that the presence of a good pressure is not a reliable sign of the absence of a clot in the catheter tip

The high incidence of this complication in our laboratory may, in part, be explained

by the fact that during the first part of this study, none of the recently suggested precautions were employed. However since the first cases occurred despite the use of all theretofore recommended modifications it is felt this type of accident may be related to the nature of the left coronary catheter (possibly the increased angulation of the catheter tip), or to the sequence in which it is used

The problem might be reduced by withdrawal and flushing of the catheter between each injection. This would however negate much of the procedural advantage (that of minimizing fluoroscopic and procedure time) and would increase the time in which a fibrin clot might form as well as predispose to other complications such as subintimal coronary dissection^{8,10} or perforation of the coronary artery¹¹

Recently, use of the guide wire and/or insertion of the second catheter has been casually implicated. Use of Teflon sheaths rather than wires and systemic anticoagulation have therefore been suggested as reasonable modifications with which to reduce the incidence of this complication⁴

At the present time we are however again limiting the transfemoral approach to only the indications previously outlined and would suggest this procedure might be less innocuous than its previous acceptance indicates

Summary

A review is presented of 139 selective coronary artery studies during a 25 month period in which 1112 coronary artery injections were performed using the percutaneous transfemoral artery approach as described by Judkins in 1967.² Six patients (four per cent) developed acute occlusion of the left coronary artery or one of its branches during the procedure and two of these died with occlusion of the left main coronary artery shortly thereafter

It is felt that since in all cases of this series and most of those of others embolic occlusion within the left coronary artery system occurred the complication is related to the thrombogenic properties of the left coronary catheter or to the procedure in which introduction of the second catheter is required

REFERENCES

- 1 Sones F M Jr and Shurey E K Cine coronary arteriography Mod Concepts Cardiovasc Dis 33:7 1962
- 2 Judkins M P Selective coronary arteriography I A percutaneous transfemoral technique Radiology 89 815 1967
- 3 Kangilaski J Medical news JAMA 221:547 1972
- 4 Spellburg R D and Ungar I The percutaneous femoral artery approach to selective coronary arteriography Circulation 36 739 1967
- 5 Gau G T Oakley C M Rahimtoola S H Raphael M J and Steiner R E Selective coronary arteriography A review of 18 months experience Clin Radiol 21 275 1970
- 6 Cheng T O Bashour I Singh B K and Kelsor G A Myocardial infarction in the absence of coronary arteriosclerosis Am J Cardiol 30 680 1972
- 7 Green G S McKinnon C M Rosch J and Judkins M P Complications of selective percutaneous transfemoral coronary arteriography and their prevention A review of 445 consecutive examinations Circulation 45 557 1972
- 8 Cheng T O Fatal thromboembolism following selective coronary arteriography Chest 62:1 1972
- 9 Wilson W J Lee G B and Amplatz H Biplane selective coronary arteriography via percutaneous transfemoral approach Am J Roentgenol 100 332 1967
- 10 Haas J M Peterson C R and Jones R C Subintimal dissection of the coronary arteries A complication of selective coronary arteriography and the transfemoral percutaneous approach Circulation 38 678 1968
- 11 Moretton L B and Wallace J M Uneventful perforation of a coronary artery during selective arteriography Am J Roentgenol 110:183 1970

Asymptomatic electrocardiographic alterations in sarcoidosis*

E Stein, MD

I Jackler, MD

B Stimmel, MD

W Stein MD

L L Siltzbach MD

New York N Y

Although sarcoidosis is not a rare disease clinical recognition of intrinsic involvement of the heart is infrequent. This is irrespective of the finding that up to 20 per cent of patients with sarcoidosis at postmortem can be demonstrated to have some degree of myocardial involvement.^{1,2} As of 1971 however only 70 cases of documented clinical involvement of the heart in sarcoidosis have been reported in the literature. The major manifestations of intrinsic cardiac involvement have been conduction disturbances, disorders of impulse formation, and progressive myocardial failure.^{1,3,6}

Recently we observed an unusual aggregation of patients with suspected cardiac sarcoidosis at the Sarcoidosis Clinic of The Mount Sinai Hospital. These patients had potentially life threatening episodes requiring hospitalization in a cardiac intensive care unit where they were continuously monitored. A review of these specific cases will be reported in detail in a separate publication.

The appearance of these serious cardiac complications prompted us to survey systematically over a three month period of

time the electrocardiographic findings of unselected asymptomatic patients seen routinely in the Sarcoidosis Clinic in an attempt to document possible asymptomatic myocardial involvement that may herald significant cardiac disease.

Patient material

In addition to the usual clinical radiologic and laboratory investigations for the presence of tissue confirmed sarcoidosis these patients were scrutinized with special attention to possible cardiac abnormalities. Patients were included in the study only if the following criteria were present: (1) under forty years of age, (2) no past or present history of cardiac or hypertensive vascular disease, (3) absence of a murmur on physical examination, (4) no history of current or past cardiotoxic or anti-hypertensive regimens, and (5) absence of cardiomegaly on x-ray. Eighty patients satisfied all of the above criteria. Twelve lead electrocardiograms (ECGs) with 30 second Lead II rhythm strips were performed on each patient using Hewlett Packard A1500 ECG machines. The ECGs were read independently by three cardi-

From the Division of Cardiology and Thoracic Diseases, Department of Medicine, Mount Sinai School of Medicine of The City University of New York, New York, N.Y.

This work was supported by a research grant from the National Heart and Lung Institute, Public Health Service (HE 13853-14).

Received for publication Dec 7, 1972.

Reprint requests to E. Stein, MD, Division of Cardiology, Mount Sinai Hospital, Fifth Ave. and 100th St., New York, N.Y. 10029.

*Read at the VIth International Conference on Sarcoidosis, Tokyo, Japan, September 11.

October 1973

Table I *Electrocardiographic alterations in 80 patients with sarcoidosis*

| | <i>Patients with abnormal ECG</i> | <i>Patients with normal ECG</i> | <i>Total patients</i> |
|---------------------------|-----------------------------------|---------------------------------|-----------------------|
| | <i>No of patients (41)</i> | <i>No of patients (39)</i> | |
| <i>Sex</i> | | | 29 |
| Male | 15 | 14 | 51 |
| Female | 26 | 25 | |
| <i>Ethnic group</i> | | | 61 |
| Black | 35 | 26 | 14 |
| Puerto Rican born | 4 | 10 | 5 |
| Caucasian | 2 | 3 | |
| <i>Estimated duration</i> | | | 33 |
| Subacute | 17 | 16 | 47 |
| Chronic | 24 | 23 | |
| <i>Chest x ray</i> | | | 32 |
| Stage I | 19 | 13 | 31 |
| Stage II | 13 | 18 | 17 |
| Stage III | 9 | 1 | |

ologists and were classified according to type of abnormality. ST segment changes were considered to be significant if altered by 1 mm or more. QT intervals were corrected for rate. The records were then reviewed with any discrepancies in interpretation subjected to discussion and final consensus.

All patients were classified as to the duration of sarcoidosis—i.e. subacute or chronic—and according to stage on the basis of chest x ray findings as described by Siltzbach.⁸

Results

There were 51 women and 29 men (Table I). 61 were black, 14 were Puerto Rican and 5 were Caucasians. Subacute sarcoidosis (of less than 2 years duration) existed in 33 patients and chronic sarcoidosis in 47 patients. Chest x rays at the time the electrocardiograms were performed showed hilar adenopathy alone (Stage I) in 32 patients, hilar adenopathy with parenchymal mottling (Stage II) in 31 patients and mottling only (Stage III) in 17 patients.

Some electrocardiographic abnormality was detected in 41 of these 80 patients without accompanying cardiac symptoms. They exhibited in all 57 separate deviations from normal. Abnormalities in rhythm and conduction were seen in 18 instances occurring in 18 patients (Table II). First de-

gree heart block was noted in 7 patients, a shortened PR interval in 5. Intra atrial conduction defects were seen in 2 patients and premature atrial contraction in one. One patient presented with AV dissociation, one with ventricular premature contractions and one with AV dissociation with junctional premature contractions.

There were also 39 repolarization abnormalities among 26 of the patients in this study (Table III). T wave changes were noted in 21 instances and consisted of flat T waves in 5, inverted in 6, notched in 9 and peaked in 1. U waves in the absence of a bradycardia were seen in 9 patients. ST segment elevation greater than 1 mm occurred in 4 patients with ST depression of equal magnitude noted in 4 others. One patient had a prolonged QT interval after correction for rate.

Factors including sex of the patient, ethnic background and duration of sarcoidosis, radiographic stage and corticosteroid therapy did not appear to influence the frequency and character of the electrocardiographic abnormalities which were found. None of the patients observed in this study exhibited clinical evidence of cardiac disease during a six month period of subsequent observation.

Discussion

In the present study greater than 50 per cent incidence of minor ECG changes in

Table II Alterations in rhythm and conduction in 15 of 80 patients with sarcoidosis

| Alterations | No of patients |
|---|----------------|
| PR intervals | |
| > 0.20 sec | 7 |
| < 0.12 sec | 5 |
| Intra atrial conduction defects | 2 |
| Premature atrial contractions | 1 |
| A-V dissociation | 1 |
| Ventricular premature contractions | 1 |
| A-V dissociation with junctional premature contractions | 1 |

Table III 39 repolarization abnormalities in 26 of 80 patients

| | No of patients |
|-------------------------|----------------|
| Flat T waves | 5 |
| Inverted T waves | 6 |
| Notched T waves | 9 |
| Peaked T waves | 1 |
| U wave (no bradycardia) | 9 |
| ST segment elevations | 4 |
| ST segment depressions | 4 |
| Prolonged QT interval | 1 |

young sarcoidosis patients was found. The significance of these alterations and its relation to clinically manifest cardiac sarcoidosis is not obvious. The nature of the underlying histopathological myocardial changes that may account for these changes in asymptomatic individuals is also unclear. The most plausible hypothesis would seem to be the fortuitous but strategic placement of sarcoid granulomas within various locations of the cardiac conducting system as simply another representation of systemic involvement in this disorder. Such a hypothesis is supported by findings of granulomas in the cardiac conducting system of patients succumbing to fatal arrhythmias.^{14,17} Whether or not myocardial granulomas have a predilection for the conducting system cannot be answered from available autopsy data or from clinical studies. Numerous granulomas are almost invariably found at autopsy in the cardiac muscle as well as in the more usual locations, suggesting that involvement of

the conduction system occurs haphazardly in a hit or miss fashion.¹⁸

The most common electrocardiographic findings reported in cases of myocardial sarcoidosis are conduction disturbances and arrhythmias consisting of frequent premature ventricular contractions, paroxysmal atrial and ventricular tachycardias, and disturbances in sinoatrial function.¹⁴ However on occasion extensive involvement of the myocardium has been seen associated with only minor ST segment and T wave changes noted on the electrocardiogram.¹¹ In one instance, the electrocardiogram in a patient with extensive myocardial sarcoidosis revealed only the presence of a sinus tachycardia.¹¹

In the literature, up to two thirds of those patients whose deaths are considered to be directly related to intrinsic cardiac sarcoidosis died suddenly.¹⁴ In 25 per cent of these fatalities no previous cardiac symptoms have been experienced. The average age at the time of death varied from 35 to 44 years.^{14,15} Porter⁴ reported fatal cardiac sarcoidosis to be twice as frequent in men as in women. However, in a series published by Mayagi and colleagues¹⁶ the frequency in the two sexes was found to be reversed. In the asymptomatic patients of our study abnormal LCG findings were equally common in both sexes.

The mechanisms leading to sudden cardiac death in sarcoidosis have not been conclusively demonstrated. However it is not unlikely that the presence of fatal arrhythmias is the common mode of exitus. It is known that individuals with abnormalities of repolarization may be more prone to develop arrhythmias through re-entry phenomena or by initiation and persistence of ectopic foci.^{16,17} Twenty six of our 80 patients demonstrated repolarization abnormalities a potentially dangerous manifestation. Fifteen patients or 18 per cent had changes in rhythm and conduction which might also represent a threat of future difficulty. The ECG alterations here reported did not correlate with the ethnic group, the duration of sarcoidosis, the severity of pulmonary sarcoidosis as determined by chest x-ray staging or treatment with corticosteroids in low dosage.

It is difficult to prove that these electrocardiographic changes are causal

of myocardial involvement without histologic evidence. However, the age of the patients studied, the absence of a past or present history of treated or untreated hypertension or heart disease and the presence of a normal sized heart on chest film make other pathogenic mechanisms less likely. In a large series of patients with proven myocardial sarcoidosis reported by Gozo and associates¹², Porter⁴ and Bashour and colleagues⁷, the electrocardiographic disturbances in rhythm and conduction which they described are similar to those seen in our series of patients.

Reversibility of cardiac involvement has been known to occur and the suggestion has been made that early corticosteroid treatment of active granulomatous disease may lead to healing of the granulomas without subsequent fibrous scarring of the myocardium.⁴ The importance of early discovery and careful follow up of cardiac involvement in sarcoidosis is evident. Early identification of patients at greatest risk may prove quite rewarding.

At the present time, prospective studies are under way in our clinic to determine which of the asymptomatic electrocardiographic abnormalities occurring in patients with sarcoidosis may represent precursors of more serious and potentially dangerous cardiac arrhythmias secondary to extensive myocardial involvement by sarcoid granulomas.

Summary

Among 80 patients with tissue confirmed sarcoidosis attending the sarcoidosis clinic of the Mount Sinai Hospital, electrocardiographic abnormalities of varying degree were noted in 41 patients despite the fact that these patients were entirely without cardiac complaints and exhibited no evidence of previous or current cardiac disease on thorough examination. All these patients were under 40 years of age.

The ECG alterations included repolarization abnormalities and alterations in rhythm and conduction. Because of the potential hazards of some of these abnormalities, prospective studies are being undertaken to determine which of them may represent precursors of dangerous intrinsic cardiac sarcoidosis.

We are grateful to Miss Maureen Baker and Miss Lynn Miller for their able technical assistance.

REFERENCES

- Longcope W and Freiman D. A study of sarcoidosis based on combined investigations of 160 cases including 30 autopsies from the Johns Hopkins Hospital and Massachusetts General Hospital. *Medicine* 31:1 1952.
- Branson J and Park J. Sarcoidosis-hepatic involvement: presentation of case with fatal liver involvement including autopsy findings and review of evidence of sarcoid involvement of liver as found in literature. *Ann Intern Med* 40:111 1954.
- Bashour F A, McConnell T, Skinner W and Hanson M. Myocardial sarcoidosis. *Dis. Chest* 88:413 1968.
- Porter C H. Sarcoid heart disease. *N Engl J Med* 263:1350 1960.
- Duvernoy W F C and Garcia R. Sarcoidosis of the heart presenting with ventricular tachycardia and atrioventricular block. *Am J Cardiol* 28:348 1971.
- Contreras R, Sanchez Torres G and Duran Rodriguez P. Sarcoidosis cardiaca. *Arch Inst Cardiol Mex* 37:20 1967.
- Shimada N, Ishihara Y, Kojima A et al. Three cases of granulomatous myocarditis with giant cells. *Acta Pathol Jap* 17:503 1967.
- Siltzbach L E. Pulmonary sarcoidosis. *Am J Surg* 55:56 1955.
- Salvesen H A. Sarcoid of Boeck: disease of importance to internal medicine: report on 4 cases. *Acta Med Scand* 86:127 1935.
- Cotter E F. Boeck's sarcoid: autopsy in case with visceral lesions. *Arch Intern Med* 64:286 1939.
- Johnson J B and Jason R S. Sarcoidosis of the heart: Report of a case and review of literature. *Am Heart J* 27:246 1944.
- Frist S M. Electrocardiographic evaluation of Boeck's sarcoid and advanced pulmonary tuberculosis: special reference to interpretation of multiple unipolar leads. *Am J Med* 7:160 1949.
- Laroche C, Gennes J L, Hazard J et al. Maladie de Besnier-Boeck-Schaumann avec manifestations polyarticulaires et localisations endomyo-pericardiques mortelles. *Bull Soc Hop Paris* 71:908 1935.
- Nissen A W and Berte J B. Cardiac arrhythmias in sarcoidosis. *Arch Intern Med* 115:275 1964.
- Miyaji T, Ohara M, Funaki M et al. A case of myocardial sarcoidosis with Adams-Stokes syndrome. *J Jap Soc Intern Med* 56:260 1967.
- Goldreyer B N. Intracardiac electrocardiography in the analysis and understanding of cardiac arrhythmias. *Ann Intern Med* 77:117 1977.
- Kistner A D and Landowne M. Retrograde conduction from premature ventricular contractions: common occurrence in the human heart. *Circulation* 37:738 1951.
- Gozo E G, Cosnow I, Cohen H and Okun L. The heart in sarcoidosis. *Chest* 60:379 1971.

Table II Alterations in rhythm and conduction in 15 of 80 patients with sarcoidosis

| Alterations | No of patients |
|---|----------------|
| PR intervals | |
| >0.20 sec | 7 |
| <0.12 sec | 5 |
| Intra atrial conduction defects | 2 |
| Premature atrial contractions | 1 |
| A-V dissociation | 1 |
| Ventricular premature contractions | 1 |
| A-V dissociation with junctional premature contractions | 1 |

Table III 39 repolarization abnormalities in 26 of 80 patients

| | No of patients |
|-------------------------|----------------|
| Flat T waves | 5 |
| Inverted T waves | 6 |
| Notched T waves | 9 |
| Peaked T waves | 1 |
| U wave (no bradycardia) | 9 |
| ST segment elevations | 4 |
| ST segment depressions | 4 |
| Prolonged QT interval | 1 |

young sarcoidosis patients was found. The significance of these alterations and its relation to clinically manifest cardiac sarcoidosis is not obvious. The nature of the underlying histopathological myocardial changes that may account for these changes in asymptomatic individuals is also unclear. The most plausible hypothesis would seem to be the fortuitous but strategic placement of sarcoid granulomas within various locations of the cardiac conducting system as simply another representation of systemic involvement in this disorder. Such a hypothesis is supported by findings of granulomas in the cardiac conducting system of patients succumbing to fatal arrhythmias.^{1,11,12} Whether or not myocardial granulomas have a predilection for the conducting system cannot be answered from available autopsy data or from clinical studies. Numerous granulomas are almost invariably found at autopsy in the cardiac muscle as well as in the more usual locations, suggesting that involvement of

the conduction system occurs haphazardly in a hit or miss fashion.¹³

The most common electrocardiographic findings reported in cases of myocardial sarcoidosis are conduction disturbances and arrhythmias consisting of frequent premature ventricular contractions, paroxysmal atrial and ventricular tachycardias, and disturbances in sinus node function.¹⁴ However, on occasion, extensive involvement of the myocardium has been seen associated with only minor ST segment and T wave changes noted on the electrocardiogram.¹⁵ In one instance, the electrocardiogram in a patient with extensive myocardial sarcoidosis revealed only the presence of a sinus tachycardia.¹⁶

In the literature up to two thirds of those patients whose deaths are considered to be directly related to intrinsic cardiac sarcoidosis died suddenly.^{1,14} In 25 per cent of these fatalities no previous cardiac symptoms have been experienced. The average age at the time of death varied from 35 to 44 years.^{1,16} Porter¹ reported fatal cardiac sarcoidosis to be twice as frequent in men as in women. However, in a series published by Miyaji and colleagues¹⁶ the frequency in the two sexes was found to be reversed. In the asymptomatic patients of our study abnormal ECG findings were equally common in both sexes.

The mechanisms leading to sudden cardiac death in sarcoidosis have not been conclusively demonstrated. However, it is not unlikely that the presence of fatal arrhythmias is the common mode of exitus. It is known that individuals with abnormalities of repolarization may be more prone to develop arrhythmias through re-entry phenomena or by initiation and persistence of ectopic foci.^{16,17} Twenty six of our 80 patients demonstrated repolarization abnormalities, a potentially dangerous manifestation. Fifteen patients or 18 per cent had changes in rhythm and conduction which might also represent a threat of future difficulty. The ECG alterations here reported did not correlate with the ethnic group, the duration of sarcoidosis, the severity of pulmonary sarcoidosis as determined by chest x-ray staging or treatment with corticosteroids in low dosage.

It is difficult to prove that these electrocardiographic changes are a result

nurse monitors stationed on medical wards use standardized self-coding sheets to record information on consecutively admitted patients. Data are collected on patient characteristics, diagnoses, the therapeutic efficacy of all drugs administered as well as full details of dosage and duration of therapy. When drug treatment is instituted, the prescribing physician is interviewed to determine the therapeutic indications. Reasons for termination of therapy and descriptions of suspected adverse reactions are recorded as well. This report is based upon data gathered since 1966 on 13 349 hospitalized medical patients in nine hospitals in the United States, Canada, Israel, and New Zealand, of whom 268 (2.0 per cent) received propranolol during one or more admissions.

Results

The mean age of the 268 propranolol recipients was 57 years and 52.2 per cent were men. Thirty-two had a discharge diagnosis of acute myocardial infarction, 19 of rheumatic heart disease, 20 of thyrotoxicosis, and 11 of paroxysmal atrial tachycardia. A wide variety of cardiovascular disorders were represented in the discharge diagnosis of the other 186 patients. Fig. 1 shows the indications for propranolol therapy. Most patients received the drug for angina (115) or cardiac arrhythmias (93), while hypertension (31) and thyrotoxicosis (20) were less frequent indications. In 4 patients the reasons for therapy were not specified.

Other cardiovascular drugs were frequently administered concurrently with propranolol. They included nitrates (nitroglycerin, isosorbide dinitrate, pentaerythritol tetranitrate) in 99 patients, digitalis glycosides in 89, diuretics (thiazides, mercurials, furosemide, ethacrynic acid, spiro lactone) in 69, antiarrhythmics (quinidine, procainamide, lidocaine, diphenylhydantoin) in 63, and antihypertensive agents (hydralazine, methyldopa, guanethidine, reserpine, thalidomide, diazoxide) in 37. Many patients received several cardiovascular drugs concomitantly.

Seven of the 268 propranolol recipients died while in the hospital, but the drug was not specifically implicated in any of these deaths.

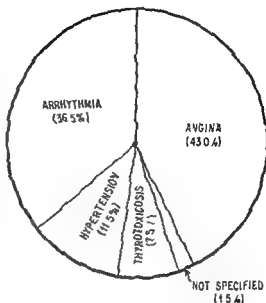


Fig. 1 Indications for propranolol in 268 patients.

Table 1 Adverse reactions to propranolol among 25 patients

| | |
|---------------------------------------|-----------|
| <i>Life threatening reactions</i> | |
| Bradycardia and shock | 3 |
| Pulmonary edema | 3 |
| Complete heart block | 1 |
| Bradycardia and angina | 1 |
| Total | 8 |
| <i>Non life threatening reactions</i> | |
| Asymptomatic bradycardia | 3 |
| Neurologic disturbances | 4 |
| Hypotension and syncope | 3 |
| Asymptomatic hypotension | 2 |
| First heart block | 1 |
| Fluid retention | 1 |
| Epigastric pain | 1 |
| Total | 17 |
| Total with adverse reactions | 25 |

Adverse reactions attributed to propranolol were reported in 25 patients (9.3 per cent). These are summarized in Table 1. Life threatening reactions occurred in 8 patients and are detailed in Table II. In four of these cases the toxic reaction occurred within hours after propranolol was first administered. In three the onset of toxicity was delayed for several days. Case 2 first developed pulmonary edema three

Adverse reactions to propranolol in hospitalized medical patients. A report from the Boston Collaborative Drug Surveillance Program*

David J Greenblatt MD**

Jan Koch Weser, MD***

Boston, Mass

Propranolol an effective beta adrenergic antagonist was introduced in 1964¹ and is now a widely prescribed drug in clinical medicine. It is extensively used either alone or in combination with other agents to treat cardiac arrhythmias of ventricular or supraventricular origin, angina pectoris, obstructive cardiomyopathies, essential hypertension and thyrotoxicosis.²⁻⁴ Propranolol may also be of benefit in some patients with hyperkinetic heart syndromes,⁵ migraine headache,⁶ essential tremor,^{7,8} anxiety,⁹ psychoses¹⁰ and septic shock.^{11,12}

Because of its ability to block beta adrenergic receptors propranolol can depress cardiac function of patients who are de-

pendent on sympathetic stimulation. For this reason it has been considered a potentially dangerous drug particularly in patients with severe heart disease or uncompensated congestive heart failure. Yet there is little information regarding the frequency of adverse reactions to propranolol or the factors predisposing to toxicity. This study presents data on adverse effects of propranolol based on observation of 268 hospitalized medical patients who received the drug.

Patients and methods

The scope and operation of the Boston Collaborative Drug Surveillance Program have been described elsewhere.^{13,16} Trained

From the Boston Collaborative Drug Surveillance Program, Boston University Medical Center, and the Clinical Pharmacology Unit, Massachusetts General Hospital, Boston, Mass. This manuscript was prepared by Drs Greenblatt and Koch Weser.

The Boston Collaborative Drug Surveillance Program is supported by Public Health Service Contract No. NIH 72 1010 from the National Institute of General Medical Sciences (NIGMS) and in part by grants from the Veterans Administration, the Canadian Food and Drug Directorate, the Israeli Ministry of Health, Auckland Hospital, Auckland, New Zealand, the Roger Williams General Hospital (Brown University), NIGMS grant No. GM 16538-02, and Hoffmann-La Roche, Inc.

Received for publication Dec. 8, 1972.

Reprint requests to: Dr. David J. Greenblatt, Boston Collaborative Drug Surveillance Program, 400 Totten Pond Rd., Waltham, Mass. 02154.

*Hospitals currently participating in the Boston Collaborative Drug Surveillance Program are: Boston, Massachusetts: Peter Bent Brigham Hospital; Boston Veterans Administration Hospital; and Massachusetts General Hospital; Providence, Rhode Island: Roger Williams General Hospital; Canada: St. Joseph's Hospital, London, Ontario; New Zealand: Auckland Hospital; Israel: Hadassah Hebrew University Hospital; NIGMS grant No. GM 16538-02, and Hoffmann-La Roche, Inc.

**Clinical and Research Fellow in Medicine and Pharmacology, Massachusetts General Hospital, supported by a Research Fellowship from the Medical Foundation, Inc., Boston, Mass.

***Chief, Clinical Pharmacology Unit, Massachusetts General Hospital; Associate Professor of Pharmacology, Harvard Medical School.

Table III Non life threatening reactions to propranolol

| Case no | Age (yrs) | Sex | Diagnosis | Indication for propranolol | Dosage | Time from initiation of therapy to onset of toxicity | Description of event |
|---------|-----------|-----|--|----------------------------|--------------------|--|--|
| 9 | 60 | F | Ischemic heart disease, diabetes mellitus, hypertension | Hypertension | 300 mg/day | 4 days | While washing hands patient complained of headache then had syncopeal episode |
| 10 | 70 | F | Ischemic heart disease hypertension PAT | Arrhythmia | 30 mg/day | 2 days | Found to be excessively drowsy and sleepy. Symptoms disappeared when drug was discontinued |
| 11 | 74 | M | Chronic glomerulonephritis, hypertension | Hypertension | 240 mg/day | Hours | Intermittent lightheadedness and dizziness concurrent with beginning therapy |
| 12 | 75 | F | Ischemic heart disease acute MI | Angina | 60 mg/day | 4 days | Bradycardia (40-50/minute) |
| 13 | 57 | M | Ischemic heart disease, acute MI | Arrhythmia | 40 mg/day | 4 days | Caused 5½ lbs. due to fluid retention |
| 14 | 78 | F | Diabetes mellitus, ischemic heart disease acute MI | Angina | 30 mg/day | 3 days | Blurring of vision |
| 15 | 50 | M | Hypertension | Hypertension | 200 mg/day | 21 days | Bradycardia (45/minute) |
| 16 | 66 | M | Ischemic heart disease | Angina | 160 mg/day | 9 days | Bradycardia (< 60/minute) |
| 17 | 15 | M | Cardiomyopathy PAT | Arrhythmia | 120 mg/day | 5 days | Hypotension (90/45) |
| 18 | 63 | M | Recurrent PAT | Arrhythmia | 30 mg/day | Hours | Bradycardia (40/minute) |
| 19 | 40 | F | Ischemic heart disease hypertension | Hypertension | 30 mg/day | 4 days | Fall in BP (210/135 to 100/90) and syncopeal episode |
| 20 | 0 | F | Diabetes mellitus, hypertension, ischemic heart disease | Angina | 30 mg/day | Hours | Fall in BP (180/110 to 100/50) and syncopeal episode |
| 21 | 36 | M | Ischemic heart disease, ventricular irritability | Arrhythmia | 20 mg/day | Hours | Bradycardia (45-50/minute) |
| 22 | 51 | F | Rheumatic heart disease atrial fibrillation congestive heart failure | Arrhythmia | 60 mg/day | 9 days | Hypotension (84/60) |
| 23 | 69 | F | Hypertension diabetes mellitus, ischemic heart disease obstructive pulmonary disease | Arrhythmia | 1 mg IV (one dose) | Immediate | 1 heart block |
| 24 | 52 | F | Diabetes mellitus, ischemic heart disease | Angina | 30 mg/day | 1 day | Nausea, headache, drowsiness, and diaphoresis |
| 25 | 60 | F | Hypertension, ischemic heart disease hiatus hernia | Angina | 40 mg/day | 3 days | Epigastric pain |

oral administration of propranolol the adverse reaction rate was independent of the dose. Two of six patients who received the drug intravenously had adverse reactions both suffering impairment of atrioventricular conduction and heart block.

Discussion

In this study interpretations of untoward clinical events as adverse reactions to propranolol were made by numer-

ous physicians in several hospitals. Rarely could the causal role of the drug be established conclusively. Nevertheless these findings suggest that the use of propranolol in hospitalized patients is associated with appreciable risks. Nine per cent of patients exposed to the drug were judged to have experienced adverse reactions.

While no deaths were directly attributed to propranolol nearly one third (8 of 25) of adverse reactions were considered life

Table III Non life threatening reactions to propranolol

| Case no | Age (yrs) | Sex | Diagnoses | Indication for propranolol | Dosage | Time from initiation of therapy to onset of toxicity | Description of event |
|---------|-----------|-----|--|----------------------------|--------------------|--|---|
| 9 | 65 | F | Ischemic heart disease diabetes mellitus, hypertension | Hypertension | 3.0 mg/day | 4 days | While washing hands patient complained of headache then had syncopal episode |
| 10 | 70 | F | Ischemic heart disease hypertension PAT | Arrhythmia | 30 mg/day | 2 days | Found to be excessively drowsy and sleepy Symptoms disappeared when drug was discontinued |
| 11 | 4 | M | Chronic glomerulonephritis hypertension | Hypertension | 40 mg/day | Hours | Intermittent lightheadedness and dizziness concurrent with beginning therapy |
| 12 | 75 | F | Ischemic heart disease acute MI | Angina | 60 mg/day | 4 days | Bradycardia (40-50/minute) |
| 13 | 57 | M | Ischemic heart disease acute MI | Arrhythmia | 40 mg/day | 4 days | Gained 5½ lbs. due to fluid retention |
| 14 | 58 | F | Diabetes mellitus, ischemic heart disease acute MI | Angina | 30 mg/day | 3 days | Blurring of vision |
| 15 | 85 | M | Hypertension | Hypertension | 400 mg/day | 21 days | Bradycardia (45/minute) |
| 16 | 55 | M | Ischemic heart disease | Angina | 160 mg/day | 9 days | Bradycardia (< 60/minute) |
| 17 | 15 | M | Cardiomyopathy PAT | Arrhythmia | 120 mg/day | 3 days | Hypotension (70/4) |
| 18 | 63 | M | Recurrent PAT | Arrhythmia | 30 mg/day | Hours | Bradycardia (40/minute) |
| 19 | 40 | F | Ischemic heart disease hypertension | Hypertension | 30 mg/day | 4 days | Fall in BP (10/135 to 1.0/90) and syncopal episode |
| 20 | 0 | F | Diabetes mellitus, hypertension, ischemic heart disease | Angina | 30 mg/day | Hours | Fall in BP (180/110 to 170/0) and syncopal episode |
| 21 | 36 | M | Ischemic heart disease ventricular irritability | Arrhythmia | 20 mg/day | Hours | Bradycardia (45-50/minute) |
| 22 | 51 | F | Rheumatic heart disease atrial fibrillation congestive heart failure | Arrhythmia | 60 mg/day | 9 days | Hypotension (84/60) |
| 23 | 69 | F | Hypertension diabetes mellitus, ischemic heart disease obstructive pulmonary disease | Arrhythmia | 1 mg IV (one dose) | Immediate | 2 I heart block |
| 24 | ■ | F | Diabetes mellitus, ischemic heart disease | Angina | 30 mg/day | 1 day | Nausea, headache drowsiness, and diaphoresis |
| 25 | ■ | F | Hypertension ischemic heart disease hiatus hernia | Angina | 40 mg/day | 3 days | Epigastric pain |

oral administration of propranolol the adverse reaction rate was independent of the dose Two of six patients who received the drug intravenously had adverse reactions both suffering impairment of atrioventricular conduction and heart block.

Discussion

In this study interpretations of untoward clinical events as adverse reactions to a drug were made by numer-

ous physicians in several hospitals Rarely could the causal role of the drug be established conclusively Nevertheless these findings suggest that the use of propranolol in hospitalized patients is associated with appreciable risks Nine per cent of patients exposed to the drug were judged to have experienced adverse reactions

While no deaths were directly attributed to propranolol nearly one third (8 of 25) of adverse reactions were considered life

Table II *Life threatening reactions to propranolol*

| Case no | Age (yrs) | Sex | Diagnoses | Indication for propranolol | Dosage | Time from initiation of therapy to onset of toxicity | Description of event |
|---------|-----------|-----|---|----------------------------|--------------------|--|---|
| 1 | 70 | F | Thyrotoxicosis atrial tachycardia | Arrhythmia | 120 mg/day | 5 days | Patient was found confused, diaphoretic, hypotensive, in slow atrial fibrillation with rate of 30-40/minute |
| 2 | 77 | M | Ischemic heart disease congestive heart failure, ventricular irritability | Arrhythmia | 30 mg/day | 3 days | Developed acute pulmonary edema requiring morphine, furosemide, hydrocortisone. Nine days later propranolol was reinstituted with immediate recurrence of pulmonary edema |
| 3 | 58 | M | Ischemic heart disease acute MI | Angina | 30 mg/day | 9 days | Acute pulmonary edema |
| 4 | 69 | M | Ischemic heart disease | Angina | 10 mg (two doses) | Hours | One hour after second 10 mg dose patient developed chest pain, hypotension, sinus bradycardia, shock |
| 5 | 63 | M | Hypertension, diabetes mellitus, ischemic heart disease, ventricular irritability | Arrhythmia | 30 mg/day | 2 days | Acute pulmonary edema |
| 6 | 55 | F | Ischemic heart disease atrial flutter | Arrhythmia | 20 mg (one dose) | 2 hours | Two hours after a single dose (20 mg po) patient developed hypotension (10/0), cyanosis, and shock |
| 7 | 76 | F | Ischemic heart disease acute MI | Arrhythmia | 1 mg IV (one dose) | Immediate | After single IV doses of digoxin (75 mg) and propranolol (1 mg) immediately developed complete heart block with slow ventricular rhythm and clinical shock |
| 8 | 79 | F | Ischemic heart disease | Arrhythmia | 10 mg (two doses) | Immediate | Bradycardia (40-60/minute) and angina shortly after the second 10 mg dose of propranolol |

days after propranolol was begun. Following nine days without the drug, the patient again received propranolol and immediately developed pulmonary edema.

Fifteen other non life threatening reactions to propranolol occurred and are detailed in Table III. Asymptomatic bradycardia occurred 5 times. In five other patients hypotension was noted, this was asymptomatic twice and associated with syncope three times. In 4 cases adverse effects involved the central nervous system; these included drowsiness, fatigue, light headedness, dizziness, headache, nausea, or blurring of vision. Second degree heart block, epigastric pain, and fluid retention each occurred once.

Although adverse reactions became more

common with increasing age, this trend did not reach statistical significance (Table IV). Reactions were significantly more common in azotemic patients (Table IV). The frequency of adverse reactions also increased significantly with the duration of hospitalization (Table IV).

A number of other factors were evaluated to determine their possible association with adverse reactions. These included sex, hospital, diagnosis, indication for propranolol, and concurrent administration of any of the previously mentioned groups of cardiovascular agents. None of these factors significantly influenced the frequency of adverse reactions to propranolol.

Finally, dosage and route of administration were considered (Table IV). During

neurotoxic manifestation occasionally reported. Hypoglycemia and bronchospasm are potential adverse effects which were not noted in this series.

The frequency of adverse reactions to propranolol was almost identical in patients receiving 100 mg per day or more and in those taking less. Seven of eight life threatening reactions occurred in patients receiving 30 mg per day or less. Equally striking is the rapidity with which many of the adverse reactions developed. Nine of the 23 toxic reactions including four of the eight life threatening events occurred within a few hours after drug therapy was initiated. These findings may reflect the fact that effective beta adrenergic blockade develops rapidly even after 10 to 20 mg doses of propranolol. They suggest that adverse hemodynamic reactions during propranolol therapy are not caused by unusually intensive beta adrenergic blockade or by a direct dose related cardiodepressive action of the drug but rather by the inability of certain patients to tolerate the usual decrease in sympathetic stimulation of the myocardium.

In patients with azotemia toxicity occurred more frequently than in those with normal renal function. In addition the adverse reaction rate in patients sixty years of age or older was higher than in those who were younger. The increase of toxicity with azotemia could be partly caused by impaired renal excretion of active metabolites of propranolol.²¹ The increase with age might be related to impaired drug metabolizing capacity in the elderly.²² However both relationships could also reflect the increased hazard of toxicity in patients with more severe underlying disease. The incidence and severity of cardiac disease may be expected to increase with age. This explanation is also suggested by the further observation that patients with long hospitalizations which presumably reflect relatively serious illnesses develop propranolol toxicity far more often than those with short hospitalizations.

Summary

Of 13 349 hospitalized medical patients monitored in a drug surveillance program 763 (2.0 per cent) received propranolol

during one or more hospitalizations. Angina and cardiac arrhythmias were the most common indications for therapy. Adverse reactions attributed to propranolol were reported in 23 patients (9.3 per cent). Fifty four reactions were considered life threatening (pulmonary edema, shock, complete heart block), all of these involved depressed myocardial function, seven occurred in patients receiving 30 mg per day or less and four occurred within four hours after propranolol was first administered. Adverse reactions were more frequent in elderly patients, in those with azotemia or long hospital stays and after intravenous administration. Propranolol dosage did not influence the frequency of adverse reactions. The findings suggest that in some patients serious adverse reactions occur rapidly even after small doses of propranolol and that patients with more severe underlying disease may be predisposed to adverse reactions.

REFERENCES

- 1 Black J W, Crowther A F, Shanks R G et al. A new adrenergic beta receptor antagonist. *Lancet* 1:1080 1964.
- 2 Fitzgerald J D. Perspectives in adrenergic beta receptor blockade. *Clin Pharmacol Ther* 10:292 1969.
- 3 Dollery C T, Paterson J W and Conolly M E. Clinical pharmacology of beta receptor blocking drugs. *Clin Pharmacol Ther* 10:765 1969.
- 4 Epstein S E and Braunwald E. Beta adrenergic receptor blocking drug. Mechanisms of action and clinical applications. *N Engl J Med* 2:5:1106 1175 1966.
- 5 Frohlich E D. Beta adrenergic blockade in the circulatory regulation of hyperkinetic states. *Am J Cardiol* 27:195 1971.
- 6 Weber R B and Reinmuth O M. The treatment of migraine with propranolol. *Neurology (Minneapolis)* 22 366 1972.
- 7 Savitt I. The effect of adrenergic beta blocking drugs on tremor. *Practitioner* 207 677 1971.
- 8 Pakkenberg H. Propranolol in essential tremor. *Lancet* 1 633 1972.
- 9 Granville-Grossman K L and Turner P. The effect of propranolol on anxiety. *Lancet* 1:788 1969.
- 10 Atsmon A, Blum I, Wiyzenbeek H et al. The short term effects of adrenergic blocking agents in a small group of psychotic patients. *Psychiatr Neurol Neurochir* 74 251 1971.
- 11 Berk J L, Hagen J F and Dunn J M. The role of beta adrenergic blockade in the treatment of septic shock. *Surg Gynecol Obstet* 130 1075 1970.

Table IV Propranolol toxicity in relation to age, renal function, and length of hospitalization

| | Number of patients | Number with adverse reaction | % |
|-------------------------------------|--------------------|------------------------------|------|
| <i>Age*</i> | | | |
| Less than 50 years | 90 | 5 | 5.6 |
| 50 to 59 years | 81 | 8 | 9.9 |
| 60 years or older | 97 | 12 | 12.4 |
| <i>Admission BUN value*</i> | | | |
| Less than 25 mg./100 ml | 200 | 13 | 6.5 |
| 25 mg./100 ml or greater | 63 | 12 | 19.0 |
| <i>Length of hospitalization***</i> | | | |
| Less than 10 days | 88 | 4 | 4.5 |
| 10 to 19 days | 124 | 10 | 8.1 |
| 20 days or more | 56 | 11 | 19.6 |

* χ^2 (2 d.f.) = 2.47 $p > .2$ ** χ^2 (1 d.f.) = 8.77 $p < .01$ *** χ^2 (2 d.f.) = 9.66 $p < .01$

Table V Propranolol toxicity in relation to dosage and route of administration

| | Number of patients | Number with adverse reaction | % |
|--------------------------------|--------------------|------------------------------|------|
| <i>Maximum daily oral dose</i> | | | |
| Less than 100 mg | 207 | 18 | 8.7 |
| 100 mg or greater | 58 | 5 | 8.8 |
| <i>Intravenous (any dose)</i> | 6 | 2 | 33.3 |
| Total | 271* | 25 | |

*Three patients received both oral and intravenous propranolol

threatening. In each of these cases the adverse reaction was related to cardiac depression manifested as slowing of heart rate, impaired A-V conduction, congestive heart failure, or hypotension. Previous reports have also stressed that serious adverse reactions to propranolol usually consist of these manifestations of cardiac depression^{2,4,17} which result from the drug's beta-adrenergic blocking activity in patients whose cardiac function is dependent upon adrenergic stimulation.

Of seventeen other reactions to propranolol which were less serious in nature, eleven also involved cardiac depression (hypotension, bradycardia, or heart block). Four adverse reactions were classified as neurologic. Increasing attention has re-

cently been focused upon the psychotropic and psychotoxic effects of beta-adrenergic blockers.¹⁸ While reduced cardiac output may be responsible for the non-specific symptoms of drowsiness, lassitude, fatigue, headache, or dizziness (as in Cases 10, 11, and 24), there is some evidence that propranolol and related drugs may produce neuropsychiatric effects unrelated to peripheral beta-blockade.¹⁸ Central sedative and anticonvulsant properties have been observed in animals. Prolonged high-dose propranolol therapy in humans has been thought to cause depression leading to suicidal attempts in a few cases.¹⁹ Insomnia, nightmares, hallucinations, and toxic psychoses have also been reported.^{17,18,20} Visual impairment (Case 14) is another probable

Experimental and laboratory reports

The relationship of coronary collateral inlet flow and retrograde flow in mongrel dogs

Anthony A. Cibulski MD BEE
Patrick H. Lehan MD
Harper A. Hellemis MD
Jackson Miss

The flow collected from a coronary artery opened to atmospheric pressure distal to its ligation (i.e. retrograde flow) was introduced by Anrep and Hausler¹ as a means to evaluate the intercoronary collateral system in the open-chest animal. Subsequently Gregg and associates² popularized the retrograde flow technique as a quantitative tool for assessing the magnitude of the intercoronary collateral system and over the last three decades the bulk of information regarding the collateral network has been uncovered with this technique. Prinzmetal and associates³ employed radioactive microsphere and erythrocyte myocardial distribution techniques to estimate the actual collateral flow delivered to the peripheral vascular bed of an occluded coronary vessel (abrupt closure) and concluded that retrograde flow underestimated the actual collateral flow. Eckstein⁴ stated that retrograde flow exceeds actual collateral flow by 10 per cent in normal dogs. This opinion was later supported by Kattus and Cregg⁵ with the reasoning that more blood would be likely to flow through the intercoronary anastomoses and out a low resistance coronary cannula opened to atmospheric pressure than through the

myocardial arterioles capillaries and veins of the ischemic myocardium. Levy Imperial and Zieske⁶ have suggested that retrograde flow is only 30 per cent of actual collateral flow on the basis of regional rubidium 86 myocardial clearance studies. Bloor and Roberts⁷ proposed that the intravascular content of isotope is responsible for falsely high myocardial rubidium 86 uptake rates (index of flow) and accounts for the disparity between the results of the retrograde flow and rubidium 86 clearance measurements. In theory the krypton 85 or xenon 133 clearance techniques⁸ for measuring myocardial flow are independent of the intravascular isotope content. In normal dogs these techniques estimate collateral flow to be approximately 25 per cent of normal (direct) perfusion rates whereas retrograde flow measurements in these animals are equivalent to 10 per cent of normal (direct) perfusion rates.⁹

Thus the relationship of retrograde flow to actual collateral flow in the mongrel dog remains controversial. In this study work was undertaken to investigate the relationship between retrograde and actual collateral flow in normal dogs and dogs subjected to chronic myocardial ischemia.

From the Department of Medicine, University of Mississippi Medical Center, Jackson, Miss.

Supported by National Institutes of Health Grant No. HE 11426.

Received for publication Sept. 18, 1972.

Reprint requests to Anthony A. Cibulski, MD, University of Mississippi Medical Center, 2500 N. State St., Jackson, Miss. 39216.

- 12 Berk J L Hagen J F Maly G et al
The treatment of shock with beta adrenergic blockade *Arch Surg* 104:46 1972
- 13 Slone D Jick H Borda I et al Drug surveillance utilizing nurse monitors An epidemiological approach, *Lancet* 2:901 1966
- 14 Borda I T Slone D and Jick H Assessment of adverse reactions within a drug surveillance program *JAMA* 205 645 1968
- 15 Slone D Gaetano L F Lipworth L et al Computer analysis of epidemiologic data on effect of drugs on hospital patients *Public Health Rep* 84:39 1969
- 16 Jick H Miettinen O S Shapiro S et al Comprehensive drug surveillance *JAMA* 213:1455 1970
- 17 Stephen S A Unwanted effects of propranolol *Am J Cardiol* 18:463 1966
- 18 Greenblatt D J and Shader R I On the psychopharmacology of beta adrenergic blockade *Curr Ther Res* 14 615 1972
- 19 Wial H J Propranolol induced depression *Br Med J* 2:50 1967
- 20 Zacharias F J Cowen K J Presti J Vickers J and Wall B G Propranolol in hypertension A study of long term therapy 1964 1970 *Am Heart J* 83:755 1972
- 21 Paterson J W Conolly M E Dollery C T et al The pharmacodynamics and metabolism of propranolol in man *Pharmacologia Clinica* 2:127 1970
- 22 O'Malley K Crooks J Duke E et al Effect of age and sex on human drug metabolism *Br Med J* 3 607 1971

with a large bore cannula for coupling to a constant pressure reservoir of easily adjustable height. The reservoir was employed to maintain the systemic pressure in experiments in which acute myocardial ischemia was induced. Following major surgical manipulations and prior to the insertion of all catheters heparin (10 mg per kilogram of body weight) was given intravenously followed by repetitive doses of 2 mg per kilogram of body weight every 30 minutes.

Retrograde flow measurements The circumflex arterial cannula was opened to atmospheric pressure (Fig 1 clamp B open clamp A fastened) and retrograde flow (RF) collected in a volumetric flask for 20 to 30 seconds and concomitantly recorded with the circumflex cannula flow probe. In chronic preparations with large retrograde flows the pressure gradient across the circumflex cannula was estimated (e.g. predetermined circumflex cannula resistance \times retrograde flow [flow probe]) and counterbalanced by adjusting the height of the retrograde flow outlet tubing to assure atmospheric to subatmospheric pressures at the arterial end of the cannula.

Regional myocardial flow measurements Regional myocardial blood flow in the circumflex (CX) and anterior descending (AD) myocardial territories was determined with the use of the krypton 85 clearance technique. A Chicago Nuclear scintillation detector collimator was positioned over the left ventricle. A two inch sodium iodide crystal was mounted 20 cm above the 10 cm diameter orifice of the collimator. Saline solutions (0.2 to 0.5 cc) of krypton 85 yielding counts of 20 000 to 50 000 per minute were delivered over periods of 5 to 10 seconds to the AD or CX myocardial regions via the modified Gregg cannula or CX arterial cannula respectively. During the period of krypton 85 infusion the CX arterial cannula system was opened to systemic pressure (Fig 1 clamp A open clamp B fastened). Under these conditions the mean intraepicardial coronary pressures of the circumflex and adjacent coronary arteries are equal (phasic differentials are unavoidable) and the collateral flow is assumed equal to zero as is the delivery of krypton 85 from one arterial cannula (e.g. the Gregg cannula) to the

adjacent coronary bed (e.g. the circumflex bed). Direct perfusion of the circumflex bed was maintained for at least 5 seconds following the termination of krypton 85 infusions. K and K_x are defined as the time constants of the logarithmic clearance curves describing desaturation of the AD and CX myocardial territories respectively. The time constants were interpreted to represent the flow per unit mass (in cubic centimeters per minute per 100 Gm) of their respective regions and were determined in the following manner: (1) The half time (e.g. $T_{1/2}$ expressed in minutes) of the linear stable phase of the krypton 85 logarithmic clearance curve was determined. $T_{1/2}$ is the duration of the interval during which the linear stable phase of the logarithmic clearance curve decreases to half of its initial clearance phase intercept value. (2) The time constant (K) was equated to $100 (0.693/T_{1/2})$ cc per minute per 100 Gm. To avoid induction of ventricular fibrillation the krypton 85 clearance phase was limited to 2 to 3 minutes in preparations subjected to severe myocardial ischemia (for example following circumflex arterial cannula occlusion in the normal dog). The initial fast phase⁸ of all clearance curves (5 to 10 seconds duration) was graphically eliminated and the stable linear phase was accepted as representative of mean clearance rates determined with infinitely long desaturation curves.¹⁰ The following six krypton 85 washout studies were performed. K and K_x were individually determined with (1) the CX cannula opened to systemic pressure (2) the CX cannula clamped and (3) the CX cannula opened to atmospheric pressure for the collection of retrograde flow. Ten minute recovery periods were allowed between the washout studies.

Change of donor coronary inlet flow This method is a resurrection of an expedient employed by Wiggers and Green¹¹ in 1936 but performed without the limitations imposed by the art of technology existing at that time. The method is illustrated in Fig 2. The increase in the inlet flow of the anterior descending artery (ΔF_A) following occlusion of the circumflex artery is interpreted as the collateral flow delivered from the anterior descending artery to

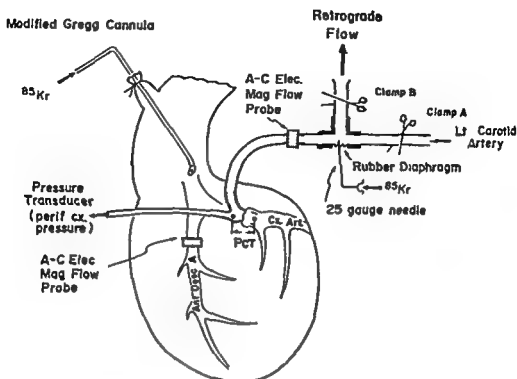


Fig 1 Perfusion of artery Ant Desc A Anterior descending artery Cx Art circumflex artery Pcr catheter pressure gradient

The approach employed was proposed to measure the total anastomotic flow delivered by the normally perfused (donor) coronary arteries to a recipient coronary artery under conditions in which the recipient vessel was either occluded or opened to atmospheric pressure for the collection of retrograde flow. The information derived with this approach was supplemented by regional flow determinations in the ischemic and nonischemic myocardial territories under conditions in which a coronary vessel was simply occluded or opened to atmospheric pressure for the collection of retrograde flow. In this study the regional myocardial blood flows were obtained with the krypton 85 clearance technique which has been shown in previous reports to provide reasonable estimates of flow under steady state conditions.⁸

Methods

The data to be reported were collected in experiments with eight mongrel dogs and ten mongrel dogs subjected to chronic myocardial ischemia by placing ameroid constrictors on their proximal circumflex arteries for periods of eight weeks or more prior to study (chronic ischemic preparations). At the time of study, all

animals (15 to 35 kilograms in weight) were anesthetized with 30 mg per kilogram of body weight of intravenously administered pentobarbital intubated and ventilated with a Harvard pump respirator. The dogs were secured in the right lateral decubitus position and a left thoracotomy was performed. The pericardium was incised and sutured to the thoracic wall to minimize movement of the heart. The proximal circumflex artery (or the segment of the vessel distal to the ameroid constrictor in the chronic preparation) was dissected, ligated proximally, cannulated just distal to the ligature and perfused from the left carotid artery through a series-coupled polyethylene tubing T tube, externally mounted A-C electromagnetic flow probe as illustrated in Fig 1. The proximal anterior descending artery was dissected free in order to implant an A-C electromagnetic flow probe. A small caliber (1 mm internal diameter) modified Gregg cannula was inserted into the left subclavian artery for transient selective catheterization of the left main stem coronary artery. The cannula was employed as a conduit for the delivery of saline solutions of krypton 85 to the left main coronary artery. The femoral artery was catheterized

Vol 86
No 4

Table 1 Krypton 85 myocardial clearance determinations

| | No of determinations | CA * cannula | | | | | |
|---|----------------------|-----------------------------|--------------|--------------|--------------|--------------------------------|--------------|
| | | Opened to systemic pressure | | Clamped | | Opened to atmospheric pressure | |
| | | \bar{A} | K_{∞} | K | K | K | \bar{A}_s |
| Normal dogs mean \pm S.D † (c.c./min./100 Gm.) | 8 | 107 \pm 24 | 110 \pm 20 | 104 \pm 22 | 24 \pm 4.5 | 110 \pm 24 | 13 \pm 2.3 |
| Chronic preparations mean \pm S.D (c.c./min./100 Gm.) | 10 | 117 \pm 19 | 105 \pm 20 | 112 \pm 20 | 97 \pm 16 | 83 \pm 22 | 21 \pm 5.8 |

K_{∞} Time constant (s/min./100 Gm.) of the krypton 85 logarithmic clearance curve describing desaturation of the anterior descending myocardium. \bar{A}_{∞} time constant (c.c./min./100 Gm.) of the krypton-85 logarithmic clearance curve describing desaturation of the circumflex myocardium.

† S.D. Standard deviation.

scending region. Following closure of the circumflex cannula the regional circumflex myocardial flow was 22 per cent ($P < 0.05$) of perfusion rates achieved with direct perfusion of the circumflex cannula under systemic pressure (i.e. normal perfusion rates). Concomitant with the collection of circumflex arterial retrograde flow the antegrade circumflex myocardial flow was determined to be 12 per cent ($P < 0.05$) of normal perfusion rates.

In chronic preparations the anterior descending myocardial flow determinations were not significantly different whether the circumflex arterial cannula was opened to systemic pressure or clamped ($K = 117$ c.c. per minute per 100 Gm. vs $K = 112$ c.c. per minute per 100 Gm. $P > 0.05$). Following the establishment of retrograde circumflex arterial flow the anterior descending myocardial flow was determined to be 71 per cent of perfusion rates measured when the circumflex arterial cannula was opened to systemic pressure ($P < 0.05$). Following occlusion of the circumflex arterial cannula the collateral flow delivered to the circumflex myocardial region represented 92 per cent of flows achieved through direct perfusion of the circumflex arterial cannula ($P > 0.05$). During the collection of retrograde circumflex arterial flow the antegrade circumflex myocardial flow persisted in magnitudes equivalent to 20 per cent of direct perfusion rates ($P < 0.05$).

During the collection of retrograde circumflex arterial flow alternative routes of krypton 85 circumflex myocardial clearance other than that resulting from effective antegrade flow in the circumflex capillary bed were determined to be insignificant. Examples follow: (1) The total blood collected retrogradely from the circumflex arterial cannula under anaerobic conditions was placed under the Chicago Nuclear collimator at a level approximated to be that of the central circumflex myocardial mass. The radioactivity in all cases (23 determinations) was less than 3 per cent of the detected decrease in radioactivity from the circumflex myocardial region during the collection of retrograde flow. The radioactivity of the retrograde flow volumes was determined to be 96 per cent (10 determinations) of an equal volume of systemic blood collected anaerobically during the period in which retrograde flow was collected. The krypton 85 content in the retrograde flow blood volumes was thus interpreted to represent krypton 85 delivered transanastomotically from the systemic circulation rather than retrograde circumflex capillary clearance. (2) At the termination of the experiment ventricular fibrillation was induced with concomitant occlusion of the anterior descending and circumflex arteries. Subsequent to the onset of ventricular fibrillation the systemic pressure was allowed to decrease spon-

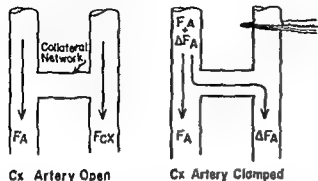
Ant Desc
Artery Circumflex
Artery

Fig 2 Donor coronary inlet flow method Following occlusion of the circumflex (Cx) artery the increase in anterior descending (Ant Desc) inflow (ΔFA) is interpreted as collateral flow delivered from the anterior descending artery to the circumflex myocardial region FA Antegrade flow to the anterior descending myocardial region FCX antegrade flow to the circumflex myocardial region delivered by direct perfusion of the circumflex artery under systemic pressure

the circumflex myocardial region providing the antegrade capillary flow to the anterior descending myocardial region remains unchanged. A similar interpretation is applied to an increase in the right coronary inflow (ΔFRC) following occlusion of the circumflex coronary artery. Thus, the total collateral flow (CF) (referred to as collateral inlet flow in subsequent discussions) to the circumflex myocardial region is

$$\text{Collateral flow (CF)} = \Delta FA + \Delta FRC$$

In this study the right coronary artery was ligated and the total collateral flow was approximated by the collateral flow delivered by the anterior descending artery.

In circumstances in which the circumflex artery is opened to atmospheric pressure for the collection of retrograde flow, the increase in flow to the donor coronary vessels is interpreted as total anastomotic flow (AF) delivered by the donor vessels to the collateral system supplying the circumflex artery.

The change in donor coronary inflow was measured 15 to 30 seconds following occlusion of the circumflex arterial cannula to allow these determinations to be made under predominantly steady state conditions. Compensatory changes in the peripheral bed of the ischemic region are essentially complete within a period of 15

seconds following occlusion of a coronary vessel in normal dogs.¹²

The donor coronary inlet flow method assumes that the flow delivered by the donor vessel (e.g., anterior descending artery) to its peripheral capillary bed remains unchanged following occlusion of the circumflex arterial cannula or opening the cannula to atmospheric pressure for the collection of retrograde flow. The constancy of the anterior descending capillary flow was evaluated by comparing the krypton 85 clearance rates (K_A) of this region prior to and following (1) circumflex cannula occlusion and (2) the collection of circumflex arterial retrograde flow.

Care was taken to employ low resistive circumflex arterial cannulas (e.g., 0.05 to 0.1 mm Hg per cubic centimeter per minute resistance) to assure accuracy of the donor coronary inlet flow method. For example, antegrade flow delivered from the left carotid artery through a high resistive circumflex cannula reduces the effective pressure head in the peripheral circumflex artery to subsystemic levels. Thus anastomotic flow delivered from donor coronary vessels may exist at a time when collateral flow is assumed to be zero (e.g., when the inlet end of the circumflex cannula is opened to systemic pressure).

The ECG and systemic pressure were monitored continuously throughout the experiment. All pressures and flows were obtained with the use of Statham strain gauge transducers and Statham alternating current electromagnetic flow probes respectively and recorded with the Electronics for Medicine recorder.

Results

Reported in Table I are the results of the regional anterior descending (K_A) and circumflex (K_C) myocardial flow determinations obtained with the krypton 85 clearance technique under the following three conditions: (1) the circumflex arterial cannula opened to systemic pressure, (2) the circumflex cannula clamped, and (3) the circumflex cannula opened to atmospheric pressure for the collection of retrograde flow.

In normal dogs there was no significant difference in the results of the three krypton 85 washout studies of the anterior de

Table 1 Krypton 85 myocardial clearance determinations

| | No of determinations | CX* cannula | | | | | |
|--|----------------------|-----------------------------|-----------------|--------------|--------------|--------------------------------|--------------|
| | | Opened to systemic pressure | | Clamped | | Opened to atmospheric pressure | |
| | | K | K _{ca} | K | K | K | K |
| Normal dogs mean \pm S.D. (c.c./min./100 Gm.) | 8 | 107 \pm 24 | 110 \pm 20 | 104 \pm 22 | 24 \pm 4.5 | 110 \pm 24 | 13 \pm 2.3 |
| Chronic preparations mean \pm S.D. (c.c./min./100 Gm.) | 10 | 117 \pm 19 | 105 \pm 20 | 112 \pm 20 | 97 \pm 16 | 83 \pm 22 | 21 \pm 3.8 |

K_{ca} Time constant (c.c./min./100 Gm.) of the krypton-85 logarithmic clearance curve describing desaturation of the anterior descending myocardium. K_{ca} time constant (c.c./min./100 Gm.) of the krypton-85 logarithmic clearance curve describing desaturation of the circumflex myocardium.
S.D. Standard deviation.

scending region. Following closure of the circumflex cannula the regional circumflex myocardial flow was 22 per cent ($P < 0.05$) of perfusion rates achieved with direct perfusion of the circumflex cannula under systemic pressure (i.e. normal perfusion rates). Concomitant with the collection of circumflex arterial retrograde flow the antegrade circumflex myocardial flow was determined to be 12 per cent ($P < 0.05$) of normal perfusion rates.

In chronic preparations the anterior descending myocardial flow determinations were not significantly different whether the circumflex arterial cannula was opened to systemic pressure or clamped ($K = 117$ c.c. per minute per 100 Gm. vs $K = 112$ c.c. per minute per 100 Gm. $P > 0.05$). Following the establishment of retrograde circumflex arterial flow the anterior descending myocardial flow was determined to be 74 per cent of perfusion rates measured when the circumflex arterial cannula was opened to systemic pressure ($P < 0.05$). Following occlusion of the circumflex arterial cannula the collateral flow delivered to the circumflex myocardial region represented 92 per cent of flows achieved through direct perfusion of the circumflex arterial cannula ($P > 0.05$). During the collection of retrograde circumflex arterial flow the antegrade circumflex myocardial flow persisted in magnitudes equivalent to 20 per cent of direct perfusion rates ($P < 0.05$).

During the collection of retrograde circumflex arterial flow alternative routes of krypton 85 circumflex myocardial clearance other than that resulting from effective antegrade flow in the circumflex capillary bed were determined to be insignificant. Examples follow: (1) The total blood collected retrogradely from the circumflex arterial cannula under anaerobic conditions was placed under the Chicago Nuclear collimator at a level approximated to be that of the central circumflex myocardial mass. The radioactivity in all cases (23 determinations) was less than 3 per cent of the detected decrease in radioactivity from the circumflex myocardial region during the collection of retrograde flow. The radioactivity of the retrograde flow volumes was determined to be 96 per cent (10 determinations) of an equal volume of systemic blood collected anaerobically during the period in which retrograde flow was collected. The krypton 85 content in the retrograde flow blood volumes was thus interpreted to represent krypton 85 delivered transanastomotically from the systemic circulation rather than retrograde circumflex capillary clearance. (2) At the termination of the experiment ventricular fibrillation was induced with concomitant occlusion of the anterior descending and circumflex arteries. Subsequent to the onset of ventricular fibrillation the systemic pressure was allowed to decrease sponta-

Table II Flow* in normal dogs

| Exp No | RF† (cc/min) | CF (cc/min) | AF (cc/min) |
|-----------------|-----------------|----------------|----------------|
| 1 | 3.7 | 5.9 | 6.5 |
| 2 | 5.0 | 10.1 | 11.6 |
| 3 | 1.6 | 4.1 | 5.1 |
| 4 | 2.9 | 7.0 | 7.8 |
| 5 | 4.0 | 8.7 | 9.7 |
| 6 | 4.7 | 9.3 | 10.7 |
| 7 | 4.5 | 9.6 | 11.1 |
| 8 | 3.2 | 8.1 | 9.8 |
| Mean \pm S.D. | 3.7 \pm 0.9 | 7.9 \pm 1.7 | 9.0 \pm 2.2 |

*Mean antegrade coronary flows A.D. 35.2 cc per minute C.V. art. 34.4 cc per minute

†RF Retrograde flow CF increase in anterior descending inflow following circumflex cannula occlusion AF increase in anterior descending inflow following the establishment of circumflex arterial retrograde flow

Table III Flow* in chronic preparations

| Exp No | RF (cc/min) | CF (cc/min) | AF (cc/min) |
|------------------|-----------------|----------------|----------------|
| 1 | 65.0 | 20.5 | 62.7 |
| 2 | 63.0 | 15.9 | 59.5 |
| 3 | 48.0 | 19.2 | 45.9 |
| 4 | 91.0 | 26.6 | 83.3 |
| 5 | 73.5 | 18.1 | 89.4 |
| 6 | 70.0 | 13.2 | 65.0 |
| 7 | 90.0 | 17.9 | 84.0 |
| 8 | 120.0 | 21.5 | 99.6 |
| 9 | 160.0 | 22.0 | 128.3 |
| 10 | 86.0 | 27.0 | 80.4 |
| Mean \pm S.D.* | 86.2 \pm 29.7 | 20.2 \pm 3.6 | 77.8 \pm 2.3 |

*Mean antegrade coronary flows A.D. = 30.5 cc per minute C.V. art. = 23.0 cc per minute

taneously and krypton 85 clearance from the myocardium was measured to be less than the equivalent of 1 cc per minute per 100 Gm (10 determinations). The clearance was assumed to be a result of evaporation. Collimator ventilation was maintained during these determinations.

The results of the donor coronary inflow and retrograde flow methods are reported for the individual experiments with normal dogs in Table II and the experiments with chronic preparations in Table III. The results of the individual experiments represents the mean of three or more determinations.

In normal dogs the retrograde flow determinations were 47 per cent ($P < 0.05$) of collateral flows (CF) estimated with the

inflow method during closure of the circumflex cannula. Retrograde flow measurements were 41 per cent ($P < 0.05$) of anastomotic flows (AF) estimated with the inflow method during the collection of retrograde circumflex arterial flow.

In chronic preparations retrograde flow measurements were 420 per cent ($P < 0.05$) of collateral flows (CF) and 111 per cent ($P > 0.05$) of anastomotic flows (AF) determined with the inflow method.

In chronic preparations the difference between the retrograde flow measurements and the anastomotic flows (AF) determined with the donor coronary inflow method during the collection of retrograde flow may be accounted for on the basis of the following mechanisms: (1) The flow to the

peripheral bed of the donor vessel decreased. This was demonstrated with the krypton 85 washout technique e.g. the anterior descending capillary flow decreased to 71 per cent of normal perfusion rates during the collection of retrograde flow (Table I). (2) Measurement of flow to the proximal anterior descending vessel does not monitor the potential contributions to anastomotic flow of the more proximal donor vessels. In the dog the primary septal artery is the major potential source of anastomotic flow proximal to the anterior descending artery. The primary septal vessel supplies approximately 10 per cent of the total left ventricular flow in the normal dog.¹¹ The contribution of the primary septal artery to retrograde flow was eliminated in five chronic preparations. In these animals the septal vessel was dissected free and acute closure of this coronary artery branch produced small discrete reductions in retrograde flow as monitored with the circumflex cannula. A C electromagnetic flow probe. The mean reduction in retrograde flow in these experiments was 4 per cent (range 1 to 7 per cent).

Discussion

In this study the hemodynamics of the coronary peripheral and collateral vascular networks were examined under conditions in which a coronary artery was simply occluded or opened to atmospheric pressure for the collection of retrograde flow. Two methods were employed to investigate the peripheral and collateral circulatory dynamics. The donor coronary inlet flow method was proposed to measure the total anastomotic flow delivered by donor coronary vessels to the coronary collateral system and the krypton 85 clearance technique was utilized to measure the regional anterior descending and circumflex myocardial blood flow.

The validity of the krypton 85 clearance technique depends upon stable flow conditions during the period of measurement. In studies in which the krypton 85 washout method was employed during a two to three minute interval following abrupt interruption of antegrade circumflex arterial flow compensatory changes in the distal circumflex bed (e.g. mainly ischemia

induced vasodilatation and decreased myocardial contractility) occur most intensely in normal dogs and primarily within the initial 10 to 15 second interval following occlusion of the circumflex cannula. Evidence for this is provided by a typical peak reactive hyperemic flow vs the duration of circumflex cannula occlusion curve.¹² In chronic preparations with well developed collaterals reactive hyperemic responses of the peripheral circumflex vascular bed were not observed following the release of temporary occlusions (> 1 minute) of the circumflex cannula. Slow changes in local coronary vasoregulation or myocardial contractility over prolonged periods (e.g. hours) were not ruled out. The purpose of this study was to measure the degree of collateral flow delivered to the circumflex bed during the initial two to three minutes following closure of the circumflex artery or the establishment of retrograde flow. In all studies the krypton 85 myocardial washout period was monitored for a minimum of two minutes thus minimizing errors resulting from early compensatory changes occurring in the distal circumflex bed. During periods of slow changes in flow the krypton 85 clearance technique provides reasonable estimates of mean flow during the interval of measurement.¹³

The krypton 85 clearance technique was employed in this study primarily to supplement rather than to compare the information derived with the donor coronary inflow method. For example the validity of the inflow method is entirely dependent upon stable flows within the peripheral donor coronary beds during periods in which the adjacent recipient myocardial region is subjected to ischemia. It was shown with the krypton 85 washout studies that the capillary flow of the anterior descending (donor) artery does not change significantly under conditions in which the circumflex cannula is clamped or opened to atmospheric pressure with the exception of the chronic preparation following the establishment of retrograde flow. In the chronic preparation the antegrade anterior descending flow decreased to 71 per cent of normal perfusion rates during the collection of retrograde circumflex arterial flow (to be discussed in a following section).

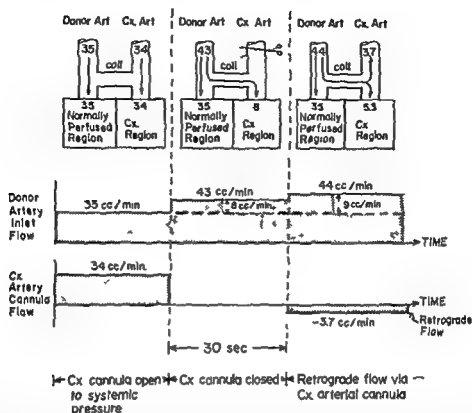


Fig 3 Circulatory dynamics. Upper drawing: Schematic representing the distribution of the donor coronary (e.g., anterior descending) inlet flow of normal dogs under conditions in which the circumflex arterial cannula is (1) open to systemic pressure, (2) clamped, and (3) opened to atmospheric pressure for the collection of retrograde flow. The numerals depicted in the upper drawing represent the mean flows (cubic centimeters per minute) reported for normal dogs in Table II. The middle and lower drawings represent the donor coronary and circumflex arterial flows (cubic centimeters per minute) respectively under the three conditions described above.

The krypton 85 washout technique yields estimates of the flow per unit mass of the myocardial region under study. Thus the donor coronary inflow and krypton 85 washout techniques are not comparable on a flow to flow basis unless the mass of the region under study is known. For this report it was chosen to compare these methods by normalizing collateral flow measurements to the normal perfusion rates (all flows are reported in Tables I to III). For example, in the normal dog, the circumflex myocardial collateral flow during closure of the circumflex artery was estimated to be 23 per cent of normal perfusion rates with the inflow method and 22 per cent of normal perfusion rates with the washout technique. The difference was not significant. In chronic preparations the collateral flow was estimated to be 88 and 92 per cent of normal perfusion rates with the inflow and washout techniques, respectively. Again the difference between the two methods was not significant.

Fig 3 is presented to provide a schematic of the peripheral and collateral circulatory dynamics which summarize the combined results of the donor coronary inlet flow and krypton 85 clearance studies in normal dogs. Illustrated in the upper portion of Fig 3 is the distribution of the donor coronary inlet flow of normal dogs (mean values are presented) under conditions in which the circumflex arterial cannula is (1) open to systemic pressure, (2) clamped, and (3) open to atmospheric pressure for the collection of retrograde flow. Following occlusion of the circumflex arterial cannula the mean donor coronary inlet flow increased from 35 to 43 cc per minute. The results of the krypton 85 clearance studies (Table I) indicate that the retrograde flow to the peripheral bed of the anterior descending artery was not significantly changed following occlusion of the circumflex arterial cannula. Therefore the increase in the donor coronary inlet flow (e.g., 8 cc per minute) is interpreted as

collateral flow delivered to the peripheral vascular bed of the occluded circumflex artery. Following the establishment of retrograde circumflex arterial flow the donor coronary inlet flow increased to 44 c.c. per minute. Again utilizing the krypton 83 clearance technique the peripheral flow of the anterior descending artery was determined to be virtually unchanged following the establishment of retrograde flow (Table I). Therefore the increase in the donor coronary inlet flow (9 c.c. per minute) is interpreted as the total anastomotic flow delivered to the collateral system during the collection of retrograde flow. Of the total transanastomotic flow (9 c.c. per minute) delivered to the recipient circumflex vascular bed 3.7 c.c. per minute are diverted retrogradely via the circumflex arterial cannula. The difference between the total anastomotic flow (9 c.c. per minute) and retrograde flow (3.7 c.c. per minute) equals 5.3 c.c. per minute and is interpreted as anastomotic flow delivered antegradely to the peripheral bed of the circumflex artery. The presence of an effective antegrade capillary perfusion of the circumflex myocardial region concomitant with the collection of retrograde circumflex arterial flow was demonstrated with krypton 83 washout studies (Table I).

In chronic preparations the flow to the peripheral bed of the anterior descending vessel determined with the krypton 83 washout method was not significantly changed following closure of the circumflex arterial cannula. Thus the increase in total donor coronary inlet flow following occlusion of the circumflex arterial cannula would reasonably be interpreted as collateral flow delivered to the recipient circumflex myocardial region. Following the establishment of retrograde circumflex arterial flow the flow to the peripheral bed of the anterior descending artery was reduced to 71 per cent of levels achieved during direct perfusion of the circumflex arterial cannula. This decreased peripheral anterior descending flow is presumably a reflection of the decreased effective perfusion pressure throughout the anterior descending epicardial network resulting from the large increase in anastomotic flow established in this bed during the collection of circumflex arterial retrograde flow. Upon

re establishment of antegrade circumflex arterial flow a marked reactive hyperemic response was noted in the circumflex arterial system and a mild reactive hyperemic response was observed in the anterior descending inlet flow. The mild reactive hyperemic response of the anterior descending inflow may be the result of one or two mechanisms. For example (1) hypoxia induced within the anterior descending bed as a result of the reduced capillary flow (i.e. 71 per cent of normal) during the collection of retrograde flow and (2) a reflection of the reactive hyperemic response in the circumflex bed. The latter mechanism is explained on the basis of the marked reactive hyperemic circumflex arterial inflows and the resultant pressure gradients generated across the circumflex cannula arterial and arteriolar systems. The reduced intra arterial pressures of the circumflex bed could thus establish transient transanastomotic pressure gradients and collateral flows temporally related to the circumflex arterial hyperemic response.

The continued perfusion of the circumflex myocardial region following the establishment of circumflex arterial retrograde flow is not unexpected since a spatial intra vascular pressure gradient must exist throughout the circumflex myocardial region falling from near systemic pressures at the collateral inlets to atmospheric pressures in the central epicardial portions of the circumflex arterial network. Thus an effective perfusion pressure is maintained for the subepicardial collateral complexes in both normal and chronic ischemic dogs during the collection of retrograde flow. In chronic preparations retrograde flow collected in the conventional manner by maintaining the circumflex arterial cannula at atmospheric pressure does not reduce the pressures in the entire circumflex epicardial bed to atmospheric levels. During the collection of retrograde flow in systems with large epicardial anastomoses an intra arterial epicardial pressure gradient must necessarily be established from the normally perfused coronary arteries to the circumflex arterial cannula. Thus the tributaries arising from at least the peripheral portions of the circumflex artery continue to receive some degree of perfusion

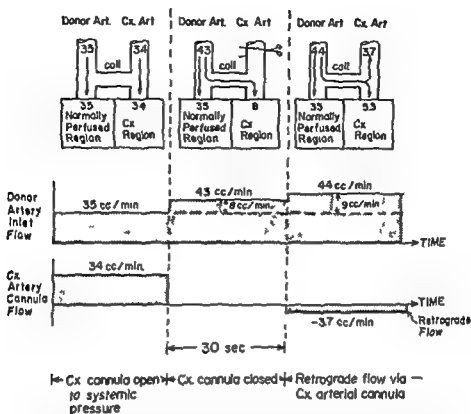


Fig. 3 Circulatory dynamics. Upper drawing, Schematic representing the distribution of the donor coronary (e.g., interior descending) inlet flow of normal dogs under conditions in which the circumflex arterial cannula is (1) open to systemic pressure (2) clamped and (3) opened to atmospheric pressure for the collection of retrograde flow. The numerals depicted in the upper drawing represent the mean flows (cubic centimeters per minute) reported for normal dogs in Table I. The middle and lower drawings represent the donor coronary and circumflex arterial flows (cubic centimeters per minute) respectively under the three conditions described above.

The krypton 85 washout technique yields estimates of the flow per unit mass of the myocardial region under study. Thus the donor coronary inflow and krypton 85 washout techniques are not comparable on a flow to flow basis unless the mass of the region under study is known. For this report it was chosen to compare these methods by normalizing collateral flow measurements to the normal perfusion rates (all flows are reported in Tables I to III). For example, in the normal dog the circumflex myocardial collateral flow during closure of the circumflex artery was estimated to be 23 per cent of normal perfusion rates with the inflow method and 22 per cent of normal perfusion rates with the washout technique. The difference was not significant. In chronic preparations the collateral flow was estimated to be 68 and 92 per cent of normal perfusion rates with the inflow and washout techniques respectively. Again the difference between the two methods was not significant.

Fig. 3 is presented to provide a schematic of the peripheral and collateral circulatory dynamics which summarize the combined results of the donor coronary inlet flow and krypton 85 clearance studies in normal dogs. Illustrated in the upper portion of Fig. 3 is the distribution of the donor coronary inlet flow of normal dogs (mean values are presented) under conditions in which the circumflex arterial cannula is (1) open to systemic pressure (2) clamped and (3) open to atmospheric pressure for the collection of retrograde flow. Following occlusion of the circumflex arterial cannula the mean donor coronary inlet flow increased from 35 to 43 cc per minute. The results of the krypton 85 clearance studies (Table I) indicate that the integrade flow to the peripheral bed of the interior descending artery was not significantly changed following occlusion of the circumflex arterial cannula. Therefore the increase in the donor coronary inlet flow (e.g. 8 cc per minute) is interpreted as

Determination of systolic intervals utilizing the carotid first derivative

Priya S Nandi MD*

David H Spodick MD**

Boston Mass

Left ventricular ejection time (LVET) is an important parameter of cardiac performance reflecting both the heart's muscle and pump functions. In one prospective blind study of unselected hospitalized patients¹ an abnormally low LVET when corrected for heart rate reliably distinguished 80 per cent of those with manifest heart disease while only 12 per cent of patients had abnormal LVET without having any manifest cardiac malfunction. This and other studies^{2,3} have shown that LVET can be used both as a reflection of functional status and as a practical screening procedure for detecting cardiac abnormality.

It has now become a standard procedure in measuring the LVET and the other systolic intervals to employ the externally recorded carotid sphygmogram. Various studies⁴⁻⁶ have proved that LVET so obtained is accurate and differs negligibly from the directly taken from the central aortic curve. However a number of technical difficulties may be encountered during the procedure of external recording of the carotid pulse. A particularly restless and uncooperative individual or persons with

obesity and respiratory problems may preclude obtaining a technically good carotid tracing despite manual expertise. Even in a cooperative patient tremors and other body movements as well as respiratory excursions may distort the base line and introduce artifacts into the carotid tracing (Fig 1). Also during exercise maintaining a stable base line of the recording may be difficult. Finally, sometimes as in aortic valvular disease the standard points of measurement—the carotid upstroke (CAR) and incisura (CAR₁)—are poorly defined even without base line or other low frequency artifacts. (This will equally invalidate the other systolic intervals.)

To obviate the above difficulties we attempted to measure the systolic intervals incorporating the first derivative of the carotid displacement pulse (dl/dt)⁶⁻¹⁰—i.e. the velocity tracing—and compare them with those obtained utilizing the carotid pulse itself. The dl/dt expressing the rate of change of the carotid displacement curve with respect to time is analogous to the rate of rise and fall of arterial pressure expressed as its first time derivative.⁶

From the Cardiology Division of the Medical Service, Massachusetts General Hospital, Boston, Massachusetts. Received for publication November 27, 1972.

Reprint requests: Dr. David H. Spodick, MD, 111 Fruit Street, Room 110, Boston, Massachusetts 02113.

**Chief of Cardiology Division, Medical Services, Lemuel Shattuck Hospital, Boston, Massachusetts. Present address: Boston University School of Medicine, Boston, Massachusetts.

*Chief of Cardiology Division, Medical Services, Lemuel Shattuck Hospital, Boston, Massachusetts. Present address: Boston University School of Medicine, Boston, Massachusetts.

during the collection of retrograde flow Iulston¹³ has demonstrated, with his stereo radiographic technique large subendo arterial collateral complexes which could provide a source of retrograde capillary flow not easily diverted by opening the circumflex arterial cannula to atmospheric pressure

The combined results of the donor coronary inflow and krypton 85 clearance studies are summarized as follows: (1) retrograde flow was 47 per cent of actual collateral flow in normal dogs; (2) retrograde flow was 420 per cent of actual collateral flow in chronic preparations reflecting the results of opening the circumflex arterial system supplied by large caliber anastomoses to atmospheric pressure and (3) concomitant with the collection of retrograde circumflex arterial flow integrative flow to the ischemic circumflex myocardium persists in both normal dogs (13 per cent of normal perfusion rates) and chronic preparations (20 per cent of normal perfusion rates)

REFERENCES

- 1 Anrep G V and Huxley H The coronary circulation I The effect of changes of the blood pressure and of the output of the heart *J Physiol* 65:357 1928
- 2 Gregg D I Thornton J J and Mautz I B The magnitude adequacy and source of the collateral blood flow and pressure in chronically occluded coronary arteries *Am J Physiol* 127:161 1939
- 3 Prinzmetal M Bergman H C Kruger H F Schwartz L I Simplicin B and Sobin S S Studies on the coronary circulation III Collateral circulation of beating human and dog hearts with coronary occlusion *AM HEART J* 35:689 1948
- 4 Eckstein R W The ineffectiveness of coronary anastomoses *Circ Res* 2:466 1954
- 5 Kuttus A A and Gregg D L Some determinants of coronary collateral blood flow in the open chest dog *Circ Res* 7:628 1959
- 6 Levy M N Imperial F S and Ziehl H Collateral blood flow to the myocardium determined by the clearance of rubidium ⁸⁶ chloride *Circ Res* 9:1035 1961
- 7 Bloor C M and Roberts L L Effects of intravascular isotope content on the isotopic determination of coronary collateral blood flow *Circ Res* 16:537 1965
- 8 Linder L Measurements of normal and collateral coronary blood flow by clo-arterial and intramyocardial injection of krypton⁸¹ and xenon¹³³ *Acta Physiol Scand* 272:5 1966
- 9 Hood Jr W B Pathophysiology of ischemic heart disease *Progr in Cardiovasc Dis* 14:297 1971
- 10 Zerler K L Equations for measuring blood flow by external monitoring of radioisotopes *Circ Res* 16:309 1965
- 11 Wiggers C J and Green H D The ineffectiveness of drugs upon collateral flow after experimental coronary occlusion in dogs *AM HEART J* 11:527 1936
- 12 Gregg D E and Fischer I C Blood supply to the heart in Hamilton W S ed *Handbook of physiology* ed 2 Washington D C 1963 American Physiological Society p 1517
- 13 Iulston W The coronary arteries Springfield Ill 1965 Charles C Thomas Publisher

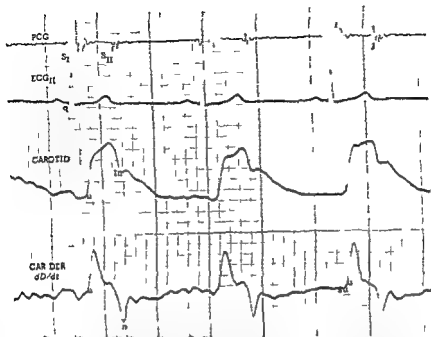


Fig 2 Top to bottom: phonocardiogram (PCG) ECG Lead II carotid displacement pulse (CAROTID) and its first time derivative pulse (CAR DER dD/dt). S_1 = First heart sound S_2 = second heart sound u = upstroke in = incisura su = slow upstroke n = nadir. Note constancy of u and variability of su .

II From carotid CAR₁ II_A

PRE-EJECTION PERIODS

- 1 From derivative (q DER) minus PTT from derivative
- II From carotid (q CAR) minus PTT from carotid

Preliminary investigation

Definition of points in the derivative

Upstroke points In the derivative trace the initial upstroke is normally slow and not always well defined when the inscription of the base line is thick or irregular. But the rapid upstroke i.e. the change of thick to thin portion of the derivative upstroke tends to be sharp and well delineated (Fig. 2). On closer inspection of the simultaneously recorded indirect carotid pulse and its first derivative it was noted that the rapid upstrokes in both the tracings were virtually simultaneous whereas the slow upstroke of the derivative was a very much earlier event (Fig. 2). We therefore included in this investigation a preliminary study comparing LVETs derived from the slow upstroke using the all ways sharp nadir of the derivative as the

end point vs. the carotid derived LVETs. Data were collected in a blinded manner as detailed earlier in 20 subjects including 10 with heart disease. It was confirmed that the mean value of the rapid upstroke of the derivative as compared to the slow upstroke was much closer to the mean value of the carotid upstroke and that derivative LVET using the rapid upstroke showed closer correlation with the carotid LVET. Results of this preliminary study are shown in Table I.

Nadir point This remains uniformly sharp and easily identifiable under all situations. Thus it serves as a very reliable measuring point when the carotid incisura may be obliterated or blunt (Fig. 1). A small but consistent difference in its timing in relation to the carotid incisura has been shown to be statistically insignificant (see below).

Results

The mean values of LVETs and other systolic intervals derived from both the carotid and its derivative traces and the comparison of the results from the two methods are shown in Tables II and III.

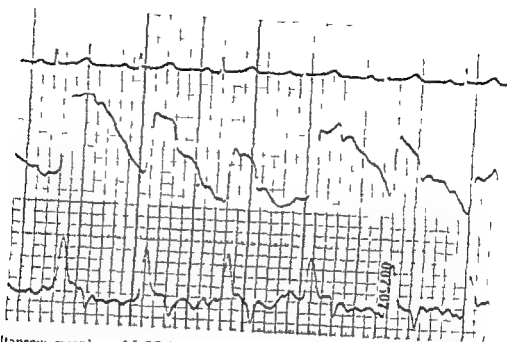


Fig. 1 Simultaneous recording of ICG Lead II indirect carotid pulse (line 2) and its first time derivative (line 3) in a patient with respiratory disease. The derivative tracing remains steady in spite of fluctuation in the carotid tracing.

Materials and method

Subjects We studied 53 ambulatory patients 30 male and 23 female aged 52 to 69. Thirty were normal subjects of whom 15 were men without clinical or electrocardiographic evidence of any heart disease the other 23 had documented hypertensive and/or coronary artery disease. These were otherwise unselected individuals drawn from a large screening clinic after exclusion only of those with arrhythmias.

Recordings With the subjects in the supine position a polygraphic recording was made as follows during relaxed expiratory apnea approximately at the same time of the day: (1) right carotid displacement pulse, using a Schwarzer photoelectric contact sensor; (2) first derivative of the carotid pulse by passing the carotid pulse wave through a RC network differentiator; technical characteristics of which are described elsewhere⁹; (3) ECG Lead II; (4) phonocardiogram at mesopex using Schwarzer contact microphone. Recording was made on a six channel Schwarzer direct writing recorder model No 622 at a paper speed of 50 mm per second with time lines at 20 msec.

Measurements The same cycles were used for both carotid and its first derivative measurements. With Q of ECG as zero time timing of the following points was measured

(Fig 2) onset of the rapid carotid upstroke (CAR_u), the carotid incisura (CAR_{in}), onset of the aortic component of the second heart sound (IIA), onset of the rapid upstroke of the first derivative (DER_u) and nadir of the derivative (DER).

Blinding procedure To avoid conscious bias measurements were made separately for the carotid and its derivative without knowledge of results obtained from the other pulse output. In each case five consecutive cardiac complexes were measured in milliseconds to a precision of 5 msec and averaged. Calculations of actual systolic intervals were made only at the end when the timings of all the above points had been measured and averaged thus avoiding any unconscious bias.

Calculations The following intervals were calculated from the measurements:

LEFT VENTRICULAR EJECTION TIME (LVET)

I From derivative

(1) Slow upstroke to nadir DER_{u-n}
 DER (used only in preliminary study)

(b) Rapid upstroke to nadir DER_u
 DER

II From carotid

Upstroke to incisura (CAR_u
 CAR_{in})

PULSE TRANSMISSION TIME (PTT)

I From derivative DER_u IIA

1.2um 36
A mbr 4

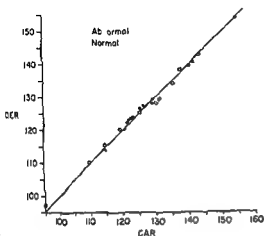


Fig 3 Relationship between the rapid upstroke points of carotid (CAR) and its derivative (DER) Statistical data in text and tables Units in milliseconds

(Rate corrections were unnecessary since measurements by each method were made in the identical cycles) The numerical values in the normal and abnormal groups were combined for final analysis to embrace larger variation and heterogeneity in clinical material

The main difference between the two upstroke points (CAR DER_a) was only 11 msec (Fig 3) which was statistically insignificant ($P > 0.5$). Thus the rapid upstrokes in both tracings may be considered to start simultaneously. This appears to be a function of the response of the RC differentiator.

The difference between the two terminal points (CAR_t DER_t) was also small (Fig 4) the DER being 5.6 msec earlier ($P > 0.2$). Thus the LVLT was statistically identical in both cases (Fig 5) the mean difference in the derivative tracing being only 4.4 msec shorter ($P > 0.2$).

The mean difference in PEP obtained from the two tracings was small (4.3 msec) and not significant ($P > 0.05$). The slightly higher P11 in the derivative trace was mainly due to the shorter P11.

P11 was significantly shorter in the derivative tracing (mean 7.7 msec, $P < 0.001$) owing to the slight but consistently early inscription (Fig 4) of the nadir point.

Discussion

In the recent years increasing attention has been paid to the development of simple

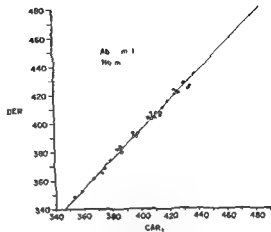


Fig 4 Relationship between the carotid incisura (CAR_t) and the derivative nadir (DER_t) points Statistical data in text and tables Units in milliseconds

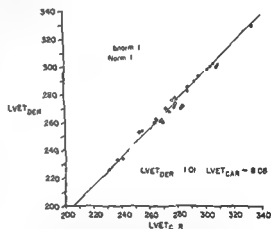


Fig 5 Relationship between two groups of LVETs derived from the carotid (LVETCAR) and its derivative (LVETDER) pulse. The regression equation is derived from the difference of slope and intercept values of the two LVETs. Units in milliseconds.

and more accurate noninvasive methods for determination of systolic time intervals. The apexcardiogram and its first derivative⁷, ear densitogram⁸ and its first derivative⁹ have all been described in these efforts. The first derivative of the carotid pulse reveals points of slope change that are often not apparent or blurred in the original curve and may thus serve as an alternate and easier noninvasive technique for this purpose.

Our results indicate that the systolic time intervals determined by utilizing the point of the carotid first derivative curve

Table I Preliminary study—LVL†*

| | Mean | SD | SE |
|--------------------|-------|-------|------|
| I Dervative | | | |
| q DER | 111.6 | 15.41 | 3.45 |
| q DER ₁ | 400.2 | 28.79 | 3.95 |
| IVET | 285.0 | 23.35 | 5.77 |
| II Carotid | | | |
| q CAR | 127.3 | 14.38 | 3.72 |
| q CAR ₁ | 404.9 | 28.69 | 6.42 |
| IVLT | 277.6 | 24.36 | 5.45 |

| | r | b | a | t | P |
|-------------------------|------|------|--------|------|-----------|
| Carotid vs dervative | | | | | |
| Upstroke points | 0.85 | 0.91 | -4.32 | 3.33 | P < 0.005 |
| Incisura or nadir point | 0.99 | 1.01 | -10.92 | 1.01 | NS† |
| LVET | 0.95 | 0.91 | 35.77 | 1.44 | NS |

*Measurement (msec) by slow upstroke (DER₁) and incisura (DER₁) of carotid first derivative and comparison with those by carotid pulse. 20 subjects.
†Not significant.

Table II Systolic intervals*

| | Mean | SD | SE |
|--------------------|-------|-------|------|
| q II _A | 479.2 | 28.09 | 3.86 |
| Carotid | | | |
| q CAR | 126.8 | 13.44 | 1.85 |
| q CAR ₁ | 405.3 | 28.13 | 3.86 |
| IVLT | 279.0 | 24.20 | 3.32 |
| PTT | 16.6 | 7.71 | 0.92 |
| PEP | 100.3 | 12.59 | 1.73 |
| Dervative | | | |
| q DER ₁ | 125.7 | 13.20 | 1.81 |
| q DER | 400.2 | 28.79 | 3.95 |
| IVET | 274.6 | 21.76 | 3.40 |
| PTT | 21.1 | 7.43 | 1.03 |
| PEP | 104.6 | 11.62 | 1.60 |

*Measurements (msec) by rapid upstroke (CAR) and incisura (CAR₁) of carotid pulse and rapid upstroke (DER₁) and nadir (DER) of carotid first derivative. 53 subjects.

Table III Comparison of systolic intervals obtained from carotid pulse and its first derivative

| | r | b | a | t | P |
|---------------------------------------|------|------|--------|------|-----------|
| 1 Upstroke points | 0.98 | 0.96 | 3.63 | 0.47 | NS |
| 2 Terminal points (incisura or nadir) | 0.99 | 1.01 | -10.92 | 1.01 | NS |
| 3 IVETs | 0.99 | 1.01 | -8.08 | 0.92 | NS |
| 4 PEPs | 0.95 | 0.87 | 16.61 | 1.82 | NS |
| 5 PTTs | 0.82 | 0.91 | -3.23 | 3.98 | P < 0.001 |

*Not significant.

Transient S-T elevation detected by 24-hour ECG monitoring during normal daily activity

Basil Golding MB ChB

Eliana Wolf MD

Dan T. noni MD

Shlomo Stern MD

Jerusalem Israel

The detection of transient S T elevation and its interpretation poses a difficult problem. That this condition has clinical significance was stressed by Prinzmetal¹ in his description of variant angina pectoris. Until recently cases with documented S T elevation were found to have this feature mainly by chance because their ECG at rest and on exercise are usually normal.² The finding of transient S T elevation during exercise in patients after myocardial infarction has been described by using radio telemetry.³ The continuous around the clock recording of the ECG by the Holter system⁴ enables the examinee to carry his normal daily routine during the monitoring and recently sporadic cases with transient S T elevation have been described using this method.^{5,6}

In this study patients in whom transient S T elevation was found by continuous ambulatory monitoring are presented their clinical background is described and the possible significance of this finding in each case is discussed.

Material and methods

The 7 patients described here were among 174 patients who were referred to us with atypical chest pain and/or palpitations for 24 hour ECG monitoring. A conventional 12 lead ECG was performed on all patients while a single Master step test and multi stage bicycle ergometric examination was performed when indicated. The Holter[®] tape recording system was used with the model E Electrocardiograph which has an improved low frequency response. Three non polarizable silver silver chloride electrodes (Mannan Creatbach) were used. The exploring electrode was placed over the V₁ position. The negative electrode was placed in the fourth intercostal space at the right sternal border and the ground electrode was placed at the manubrium sterni. The complex obtained closely resembled Lead V₁ of the conventional ECG. The tapes were screened for possible ECG abnormalities on the Electrocardioscanner and the suspicious segments seen on the screen were printed out on regular ECG paper.

From the Cardiology Service and Department of Physiological Medicine and Rehabilitation and Medicine, A Hadassah University Hospital, Jerusalem, Israel. Address reprint requests to Dr. Golding at the Hadassah Medical School, Jerusalem, Israel.
Received by the J. S. F. of the H. B. W. L. City of the Hadassah Medical Organization, IM, N. G. on, S. L. (Land).
Received for publication Dec 6, 1972.
Revised version received May 2, 1973. Address reprint requests to Dr. Golding at the Hadassah University Hospital, P.O. Box 499, Jerusalem, Israel.
Address reprint requests to Dr. Golding at the Hadassah Medical School, Jerusalem, Israel.

correlated very well with those obtained using the carotid displacement pulse itself both in normal subjects and in those with heart disease. There was no statistically significant difference between the mean values of ejection times obtained by either method. This remained true whether the ejection time is calculated from the slow upstroke or the rapid upstroke points of the first derivative, although the LVET derived from the latter point correlated more closely (i.e. slope value nearer to 1 and intercept closer to 0) with the carotid LVET.

Both the upstroke and nadir points in the derivative curves were earlier than those in the carotid tracing. This was not entirely unanticipated since there is some phase shift to be expected in the derivative due to the equivalent of a shorter time constant as a characteristic of the RC network differentiator. That the points in the time derivative trace appeared to be more reliable is also understandable since both the rapid carotid upstroke and the minimally represent the points of change of slope and these may be poorly defined in carotid curves owing to technical factors and low frequency artifacts which do not appear in time derivative curves.

The consistently early inscription of the derivative nadir with respect to the carotid measure may be fortuitous. Normally the measure of the central aortic pulse is registered with a slight delay after the aortic valve closure, perhaps due to the continued small forward flow during early left ventricular relaxation. Kumar and Luisada¹⁰ found this delay to be around 17.6 msec. In this study, the nadir of the derivative in relation to the carotid measure, has been early by an average of 7.6 msec. Thus the derivative nadir may accidentally reflect better than the carotid measure the time of the actual change of slope of the central pulse and therefore a consistent underestimation of LVET thus obtained may be really closer to the actual ejection period.

Summary

Left ventricular ejection time (LVET) is a useful parameter of cardiac function

which may be difficult to measure in rest, less, dyspneic and uncooperative patients. Certain well defined points of the first time derivative of the carotid displacement pulse are suitable to determine LVET and appear more reliable than the corresponding points in the carotid pulse itself under technically adverse situations. Systolic time intervals thus obtained correlate well with those determined by utilizing the undifferentiated carotid pulse wave. Thus the carotid first time derivative (dD/dt) serves as a more versatile and convenient method for such measurements because of greater freedom from motion artifact and better definition of points.

REFERENCES

- 1 Spodick D H, Dorr C A and Calabrese B F. Detection of cardiac abnormality by clinical measurements of left ventricular ejection time. *JAMA* 209:239 1969.
- 2 Weisler A M, Peeler R G and Roehll W H Jr. Relationships between left ventricular ejection time, stroke volume and heart rate in normal individuals and patients with cardiovascular disease. *Am Heart J* 62:367 1961.
- 3 Shalhoub H R, Sinno M Z, Chuganina R, Loeb H S, Rosen K M and Gunnar R M. Effects of occlusion on left ventricle in acute myocardial infarction. *N Engl J Med* 287:527 1972.
- 4 Shaver J A, Kroetz F W, Leonard J I and Poley H W. The effect of steady rate increase in systemic arterial pressure on the duration of left ventricular ejection time. *J Clin Invest* 37:217 1968.
- 5 Willemis J and Kesteloot H. The left ventricular ejection time. Its relation to heart rate, mechanical systole and some anthropometric data. *Acta Cardiol* 22:401 1967.
- 6 Khan A H and Spodick D H. The first derivative of the carotid displacement pulse. *Am Heart J* 81:470 1972.
- 7 Gabor G, Torubaszky I and Kalman P. Determination of systolic time intervals using the apexcardiogram and its first derivative. *Am J Cardiol* 30:217 1972.
- 8 Christie R and Spodick D H. Densitography: A new method for evaluation of cardiac performance at rest and during exercise. *Am Heart J* 83:493 1972.
- 9 Pigott V M, Christie R and Spodick D H. New methods for noninvasive determination of left ventricular ejection time during cardiovascular challenges. *Proc 25th Annual Conf Engin Med Biol (ACFMB)* 14:183 1972.
- 10 Kumar S and Luisada A A. Mechanism of changes in the second heart sound in aortic stenosis. *Am J Cardiol* 28:162 1971.

| Reason for monitoring | Findings during monitoring | Diagnosis |
|-----------------------------|---|-------------------------------------|
| Palpitations | Abrupt changes in rate from 84 to 140 inversion of T waves S-T elevation of 2 mm | Neurocirculatory asthenia suspected |
| Chest pain | Repeated S-T elevation of 2 mm T wave inversion | Neurocirculatory asthenia |
| Chest pain dizzy spells | S-T elevation of 2 mm S-T depression of 1 mm supraventricular tachycardia ventricular premature beats | Post myocardial infarction |
| Chest pain and palpitations | S-T elevation of 1.5 mm S-T depression and T wave inversion atrial premature beats and atrial tachycardia | Post myocardial infarction |
| Chest pain palpitations | S-T elevation of 2 mm | Post myocardial infarction |
| Chest pain breathlessness | S-T elevation of 3.4 mm | Variant angina pectoris |
| Chest pain | S-T elevation of 4.5 mm | Variant angina pectoris |

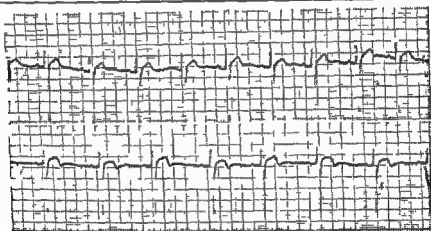


Fig 2 Upper panel ECG tracing of patient No 1 as recorded during most of the 24 hour monitoring period Lower panel S-T elevation observed during sleep lasting for 5 minutes and not causing him to awake

strip with ST elevation showed QRS complexes identical or closely resembling for direction and amplitude those of the preceding strip without the ST elevation and the amplitude of the QRS closely approximated the amplitude of V_1 in the standard ECG

Results

The seven patients who showed transient ST elevation on 24 hour monitoring were invariably male and their age varied from 18 to 66 years (Table 1)

In view of their clinical background and subsequent ECG findings the patients were divided into three groups

1 Patients with neurocirculatory asthenia

CASE 1 B Y 18 year old male referred because of palpitations which troubled him at rest and during exercise Physical examination was negative The resting ECG showed a single atrial premature beat A Master test showed no ST T changes but several atrial premature beats were recorded some of which were blocked On ergometric examination¹⁰ at a load of 100 Watts per minute and at a heart rate of 160 per minute inverted T waves in I and flat T waves in V_1 were noted The 24 hour monitoring during the day detected sinus tachycardia of 170 per minute when the patient ascended a flight of stairs or performed other relatively light physical effort and this was usually accompanied by T wave inversion S-T elevation of 2 mm was recorded during sleep lasting for 5 minutes and not causing him to awake (Fig 2) A presumptive diagnosis of neurocirculatory asthenia was made in this patient.

Table 1 Details of 7 patients with transient ST elevation detected by continuous 24 hour monitoring

| Patient No and initials | Age (yr) | Sex | Resting ECG | Master two step test | Bicycle exercise test |
|-------------------------|----------|-----|---|------------------------------|---|
| 1 B Y | 18 | M | Normal | Normal | Normal |
| 2 C A | 28 | M | T wave inversion | T wave inversion more marked | — |
| 3 F C | 61 | M | Antero-septal infarction | — | S-T depression of 1 mm in V_4 and ventricular premature beats |
| 4 H A | 56 | M | Antero-septal infarction | Positive | S-T depression of 3 mm in V_4 |
| 5 M B | 54 | M | Anteroseptal infarction | — | — |
| 6 H Y | 66 | M | Right B B B ventricular premature beats | — | — |
| 7 B Z | 47 | M | Right B B B | Normal | — |



Fig 1 Step voltage decay curve of a Sanborn 100 ECG machine (solid line) and of the Holter Avionics system (interrupted line). It can be seen that after 200 msec both systems have similar decay. A 90 per cent decay is reached in the Sanborn 100 machine after 300 msec and in the Holter Avionics system after 240 msec.

using an improved method of operation.⁸ The reliability of the system is to reproduction of ST segment was tested earlier by comparing the tracings of individuals with normal ECG pattern and of others with pathological ST T changes recorded through a conventional ECG machine with that obtained through the Avionics Electrocardiometer E and its interpreting equipment the results demonstrated identity of the tracings.⁸ Furthermore the Electrocardiometer and reproduction system have a correction network for improving its low frequency response which is not a simple RC network. Therefore we compared the low frequency response of a conventional ECG machine (Sanborn 100) and the Electrocardiometer E by applying to both of them a step voltage. Fig 1 shows that the step voltage decays to 90 per cent of its original value in the conventional

ECG machine after 300 msec and in the Avionics system after 240 msec. Since the duration of the ST segment is not longer than 200 msec and after this interval the decay in the step voltage in the Avionics system is similar to that of the conventional ECG (92 per cent and 93 per cent respectively) we conclude that for clinical purposes this system is satisfactory for reproducing the ST segment.

The patients were requested to keep a record of their activities and symptoms while being taped so that these could be subsequently correlated with ECG findings.

ST elevation was regarded as significant if the following conditions were fulfilled: it occurred for at least 12 seconds; the ECG strip had a stable isoelectric line (defined as being a straight line drawn through three subsequent TP segments); the ST segment being 1.5 mm or more above the baseline.

mental obstruction of 75 per cent in his right main coronary artery. We believe that with the development of selective coronary arteriography and bypass surgery the exact diagnosis of a condition with the above mentioned pathological associations and a poor prognosis has special importance. Since the history in this syndrome is atypical and both the resting and the exercise ECG are usually normal continuous ECG monitoring under dynamic conditions as described above is of particular assistance in diagnosing these cases.

Summary

Seven out of 174 patients subjected to continuous 24 hour ECG monitoring because of atypical chest pain and/or palpitations were found to have transient ST elevation. Two were diagnosed as suffering from neurocirculatory asthenia, three had myocardial infarction and two suffered most probably from Prinzmetal's variant angina pectoris. We believe that the finding of transient ST elevation is of special significance because of the pathological and therapeutic implications and that continuous ECG monitoring during unrestricted daily activities provides a unique opportunity to detect this condition more frequently.

We are greatly indebted to Prof. J. Weismann of the Bioengineering Dept. of the Hebrew University-Hadassah Medical School and to Dr. Y. Mahler of the Electronics Dept. of the Hadassah University Hospital for their useful help. The devoted technical assistance of Mrs. Batia Glassman is gratefully acknowledged.

REFERENCES

1. Prinzmetal M, Krennauer R, Wada T and Bor N. Angina pectoris I. A variant form of

- angina pectoris. Preliminary report. *Am J Med* 27:375 1959.
2. Peretz D. I. Variant angina pectoris of Prinzmetal. *Can Med Assoc. J* 80:110 1961.
3. Bellet S, Eliahu M, Deliyannis S and La Van D. Radioelectrocardiography during exercise in patients with angina pectoris compared with the postexercise electrocardiogram. *Circulation* 29: 5 1962.
4. Gilson J, Holter N Y and Glasscock W II. Clinical observations using the electrocardiogram ALSEP continuous electrocardiographic system. Tentative standards and typical patterns. *Am J Cardiol* 14: 204 1964.
5. Lunger M and Shapiro A. Continuous electrocardiographic monitoring in nocturnal angina. (Abstr.) *Am J Cardiol* 13:119 1964.
6. Whiting R B, Klein M D, Vander Veer J and Lowy B. Variant angina pectoris. *N Engl J Med* 282: 709 1970.
7. Silverman M and Flamm M D. Variant angina pectoris: anatomic findings and prognostic implications. *Ann Intern Med* 75:339 1971.
8. Tavoni D and Stern S. Improved method of operating the Holter Avionics ECG recording system. *AM HEART J* 83: 846 1972.
9. Stern S and Tavoni D. The reliability of the Holter Avionics system in reproducing the ST-T segment. (Letter to the Editor). *AM HEART J* 81: 427 1972.
10. Konig H and Messin R. Methods of evaluation of the physical capacity. *Acta Cardiol (Suppl)* 14: 30 1970.
11. Holmgren A, Jonsson M, Levander M, Linderholm H, Sjustrand T and Strom G. ECG changes in vasoregulatory asthenia and the effect of physical training. *Acta Med Scand* 162: 259 1959.
12. Frohlich M D, Dustan H P and Page I H. Hyperdynamic beta adrenergic regulatory state. *Arch Intern Med* 117: 614 1966.
13. Friedberg C K. Diseases of the heart. Philadelphia and London 1966. W B Saunders Company. p. 710.
14. Macaluso R N and Krattus A A. Angina pectoris at rest with preservation of exercise capacity—Angina inversa. (Abstr.) *Circulation (Suppl II)* 176 1967.

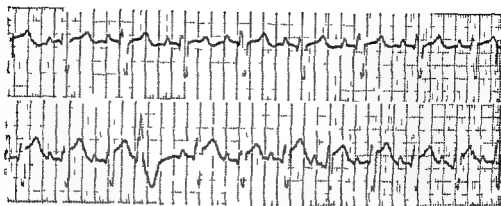


Fig 7 Upper panel LCG tracing of patient No 6 as recorded during, most of the 24 hour monitoring period. Lower panel ST elevation recorded during walking accompanied by chest pain.

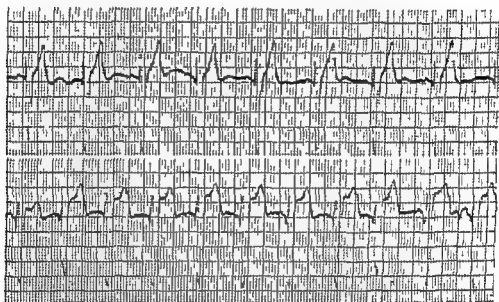


Fig 8 Upper panel LCG tracing of patient No 7 as recorded during, most of the 24 hour monitoring period. Lower panel ST elevation recorded during, a 7 minute period of chest pain.

therapy.¹² The detection of ST elevation in the two cases described above suggests the possibility of sympathetic induced coronary artery constriction. Since our patients were relatively young we assume that they have normal coronary arteries however that vasospasm in a normal artery can result in ST elevation remains to be proved.

The second group of patients consisted of three who had suffered a previous myocardial infarction. It is known that acute ischemia in cardiac muscle alongside an infarcted area may give rise to ST elevation¹³ and it is thought to explain this finding in the patients of this group.

The third group had atypical chest pain and ST elevation at rest or during ordinary physical activity. All these findings are

compatible with the diagnosis of Prinzmetal variant angina pectoris.¹ This condition implies the presence of a high grade localized occlusion in the proximal part of one of the three main coronary arteries. Prinzmetal and co workers¹ mention three such cases. Peretz and Silverman and Hamm⁷ describe one case each who at post mortem examination show severe localized stenosis at the proximal end of one of the three main coronary arteries. Macalpin and Krattus¹⁴ performed coronary arteriography on six cases and found a single major stenotic lesion in five. They suggested the possibility of surgery in these patients which would seem reasonable on the basis of the poor prognosis in this syndrome.¹ This step was taken by Silverman and Hamm⁷ in a patient who showed a seg-

mental obstruction of 75 per cent in his right main coronary artery. We believe that with the development of selective coronary arteriography and bypass surgery the exact diagnosis of a condition with the above mentioned pathological associations and a poor prognosis has special importance. Since the history in this syndrome is atypical and both the resting and the exercise ECG are usually normal continuous ECG monitoring under dynamic conditions as described above is of particular assistance in diagnosing these cases.

Summary

Seven out of 174 patients subjected to continuous 24 hour ECG monitoring because of atypical chest pain and/or palpitations were found to have transient ST elevation. Two were diagnosed as suffering from neurocirculatory asthenia, three had myocardial infarction and two suffered most probably, from Prinzmetal's variant angina pectoris. We believe that the finding of transient ST elevation is of special significance because of the pathological and therapeutic implications and that continuous ECG monitoring during unrestricted daily activities provides a unique opportunity to detect this condition more frequently.

We are greatly indebted to Prof. J. Weismann of the Bioengineering Dept. of the Hebrew University, Hadassah Medical School and to Dr. Y. Mahler of the Electronics Dept. of the Hadassah University Hospital for their useful help. The devoted technical assistance of Mrs. Batta Glasman is gratefully acknowledged.

REFERENCES

1. Prinzmetal M, Kennamer R, Wada T and Bor N. Angina pectoris I. A variant form of

- angina pectoris. Preliminary report. *Am J Med* 27:375 1959.
2. Peretz D I. Variant angina pectoris of Prinzmetal. *Can Med Assoc J* 88:110 1961.
3. Bellet S, Eshakim M, Deliyannis S and La Van D. Radioelectrocardiography during exercise in patients with angina pectoris compared with the postexercise electrocardiogram. *Circulation* 20:5 1962.
4. Gilson J, Holter N Y and Glascock W R. Clinical observation using the electrocardiographic AVSEP continuous electrocardiographic system. Tentative standards and typical patterns. *Am J Cardiol* 14:204 1964.
5. Langer M and Shapiro A. Continuous electrocardiographic monitoring in nocturnal angina (Abstr.). *Am J Cardiol* 13:119 1964.
6. Whiting R B, Klein M D, Vander Veer J and Lown B. Variant angina pectoris. *N Engl J Med* 282:109 1970.
7. Silverman M and Flamm M D. Variant angina pectoris: anatomic findings and prognostic implications. *Ann Intern Med* 75:339 1971.
8. Tzivoni D and Stern S. Improved method of operating the Holter Avionics ECG recording system. *Am Heart J* 83:846 1972.
9. Stern S and Tzivoni D. The reliability of the Holter Avionics system in reproducing the ST-T segment. (Letter to the Editor). *Am Heart J* 84:427 1972.
10. Kong K and Messin R. Methods of evaluation of the physical capacity. *Acta Cardiol (Suppl)* 14:30 1970.
11. Holmgren A, Jonsen B, Leyander M, Linderholm H, Sjostrom T and Strom G. ECG changes in vasoregulatory asthenia and the effect of physical training. *Acta Med Scand* 164:259 1959.
12. Frohlich E D, Dustan H I and Page I H. Hyperdynamic beta adrenergic circulatory state. *Arch Intern Med* 117:614 1966.
13. Friedberg C K. Diseases of the heart. Philadelphia and London 1966. W B Saunders Company p 710.
14. Vaccapio R N and Krattus A A. Angina pectoris at rest with preservation of exercise capacity—Angina inversa (Abstr.). *Circulation (Suppl 1)* 176 1967.

Depression of cardiac performance by ethanol unmasked during autonomic blockade

Maylene Wong M D *
Los Angeles, Calif

The published accounts of the hemodynamic effects of acutely administered ethyl alcohol on the cardiovascular system appear to be inconsistent¹ The literature suggests that alcohol can both stimulate and depress the heart and in some instances produce no discernible change in hemodynamics Plausible explanations for the variance are found in the differences among subjects and their preparation, in the dose of alcohol administered and in the methods and timing of measurements Conducting experiments in identical fashion in anesthetized dogs we unexpectedly made all three observations of stimulation depression and apparent unresponsiveness Consequently it seemed to us that the observations might be valid and that another explanation for the inconsistency of the findings was wanting Surmising that extracardiac stimuli especially catecholamine release by alcohol were obscuring alcohol's direct effect on cardiac performance studies were combined with autonomic blockade It is the purpose of this paper to present the data from these experiments and to offer a concept that makes the literature more consistent

Methods

Twenty mongrel dogs weighing 20 to 31.7 kilograms were studied in the fasting state All of the animals had been housed at our animal facility for at least three weeks before the experiment and had been fed with vitamin supplements once a day Their general nutrition appeared to be good and their hematocrits ranged from 39 to 55 per cent Morphine sulfate 1.0 mg per kilogram of body weight was administered subcutaneously and anesthesia was induced one hour later with chloralose 60 mg and urethane 600 mg per kilogram of body weight given intravenously Thereafter $\frac{1}{2}$ to $\frac{1}{4}$ of the anesthetizing dose was added every 60 to 90 minutes Ventilation with 100 per cent oxygen was assisted through a cuffed endotracheal tube and the volume and rate of the respirator were regulated to maintain arterial PCO_2 between 35 and 45 mm Hg Hyperinflation to 25 cm H₂O was applied automatically every 5 minutes to minimize atelectasis The electrocardiogram was monitored continuously and body temperature was recorded periodically from a thermistor placed in the ascending aorta Catheters

From the Medical Service Veterans Administration Wadsworth Hospital Center Los Angeles Calif and The Department of Medicine University of California at Los Angeles School of Medicine Los Angeles Calif
This work was supported by a Los Angeles County Heart Association Grant and No 452 and Institutional Research Funds

Received for publication Dec 18 1972

Reprint requests to Maylene Wong M D Cardiology Section Wadsworth Hospital Center Los Angeles Calif 90073

*Assistant Professor of Medicine in Residence University of California at Los Angeles School of Medicine

5 / me 86
1 mb 4

were inserted through arterial and venous cutdowns into the left ventricle, thoracic aorta, pulmonary artery and inferior vena cava and were kept patent with periodic flushes of 4 per cent sodium citrate. Aortic and left ventricular pressures were detected with Statham P23D strain gauges and zero reference for pressures was set at the midchest level. Left ventricular end diastolic pressure was recorded at the highest amplifier gain and was read during end expiration. The frequency response characteristics of the left ventricular catheter showed resonant frequencies between 30 to 40 cycles per second. Cardiac output was estimated from duplicate indocyanine green dilution curves. One ml of dye was flushed into either the pulmonary artery or left ventricle and blood was sampled at 38.2 ml per minute from the thoracic aorta through a Gilford densitometer. The blood sample was reinfused after the inscription of each curve. Calibration of the indicator was accomplished through the same catheter system using the animal's blood with the initial hematocrit and three dilutions of dye. Hematocrits at the end of the experiments were always higher than those at the start by an average of four percent age points. The difference between duplicate determinations of cardiac output was 3.3 per cent ± 1.2 (mean \pm S.E.M.).

Combined beta adrenergic and postganglionic parasympathetic blockade was produced by an intravenous injection of 0.2 mg per kilogram of body weight of propranolol (dl propranolol) and 1.0 mg per kilogram of body weight of atropine and maintained by a constant infusion of propranolol 0.004 and of atropine 0.02 mg per kilogram of body weight per minute respectively.² It has been shown in six anesthetized and atropinized dogs that after this initial dose of propranolol myocardial contractile force and heart rate do not respond significantly to stimulation of the left stellate ganglion (2 millisecond pulses at supramaximal voltage of 20 per second) or to intravenous injections of isoproterenol (0.4 mg per kilogram of body weight) for 20 minutes or more.³

The animals were grouped into 8 dogs that received infusions of ethanol with autonomic blockade (E+B), 4 dogs that

received infusions of saline with blockade (S+B) and 4 dogs that received ethanol without blockade (E-B). Six periods of observation were made in each experiment: 2 control periods at 3 and 3½ hours after anesthesia which were also 30 and 60 minutes after autonomic blockade in the blockade experiments and 4 serial observations at 30, 60, 90 and 120 minutes after starting the infusions of ethanol or saline. The infusion rates of alcohol ranged from 25 to 33 mg per kilogram of body weight per minute and averaged 29 mg per kilogram of body weight per minute for both ethanol groups and produced blood levels of ethanol over 100 mg per 100 milliliters by 30 minutes and over 200 mg per 100 milliliters by 2 hours.⁴ The concentration of ethanol was adjusted so that the volume was delivered at 2 ml per minute the same rate at which saline was infused.

In 5 of the 8 dogs receiving ethanol with blockade and 2 of the 4 animals receiving saline with blockade, 3 to 4 endocardial biopsies were sampled from the right ventricular endocardium during the experiment at spaced intervals for biochemical analyses that will be reported separately. In our hands this procedure had no measurable effect on the hemodynamics judging from negligible changes measured in a group of 4 biopsied animals comprised of the 2 from the present saline with blockade group and 2 others that received glucose with blockade. Mean changes from control measurements at the end of the 1½ hour observation period were: mean aortic pressure = -2.0 per cent, cardiac output = $+1.0$ per cent, heart rate = $+4.0$ per cent and left ventricular end diastolic pressure = $+0.1$ mm Hg.

Results

Table 1 gives the mean values for cardiac index (CI), mean aortic pressure (MAP), left ventricular end-diastolic pressure (LVEDP) and heart rate for each group at control and experimental periods. The only significant differences at the 5 per cent level of confidence between groups during the control period were a lower MAP in the ethanol with blockade (E+B) group and a slower heart rate in the ethanol without blockade (E-B) group. The com-

Depression of cardiac performance by ethanol unmasked during autonomic blockade

Maylene Wong, MD*
Los Angeles Calif

The published accounts of the hemodynamic effects of acutely administered ethyl alcohol on the cardiovascular system appear to be inconsistent.¹ The literature suggests that alcohol can both stimulate and depress the heart and in some instances, produce no discernible change in hemodynamics. Plausible explanations for the variance are found in the differences among subjects and their preparation in the dose of alcohol administered and in the methods and timing of measurements. Conducting experiments in identical fashion in anesthetized dogs we unexpectedly made all three observations of stimulation, depression and apparent unresponsiveness. Consequently, it seemed to us that the observations might be valid and that another explanation for the inconsistency of the findings was wanting. Surmising that extracardiac stimuli, especially catecholamine release by alcohol, were obscuring alcohol's direct effect on cardiac performance, studies were combined with autonomic blockade. It is the purpose of this paper to present the data from these experiments and to offer a concept that makes the literature more consistent.

Methods

Twenty mongrel dogs weighing 20 to 31.7 kilograms were studied in the fasting state. All of the animals had been housed at our animal facility for at least three weeks before the experiment and had been fed with vitamin supplements once a day. Their general nutrition appeared to be good and their hematocrits ranged from 39 to 55 per cent. Morphine sulfate 10 mg per kilogram of body weight was administered subcutaneously and anesthesia was induced one hour later with chloralose 60 mg and urethane 600 mg per kilogram of body weight given intravenously. Thereafter, 1/4 to 1/2 of the anesthetizing dose was added every 60 to 90 minutes. Ventilation with 100 per cent oxygen was assisted through a cuffed endotracheal tube and the volume and rate of the respirator were regulated to maintain arterial PCO₂ between 35 and 45 mm Hg. Hyperinflation to 25 cm H₂O was applied automatically every 5 minutes to minimize atelectasis. The electrocardiogram was monitored continuously and body temperature was recorded periodically from a thermistor placed in the ascending aorta. Catheters

From the Medical Service, Veterans Administration Woodworth Hospital Center, Los Angeles, Calif. and The Department of Medicine, University of California at Los Angeles School of Medicine, Los Angeles, Calif.
This work was supported by a Los Angeles County Heart Association Grant in Aid No. 452 and Institutional Research Funds.

Received for publication Dec. 18, 1972.

Reprint requests to Maylene Wong, MD, Cardiology Section, Woodworth Hospital Center, Los Angeles, Calif. 90033.

*Assistant Professor of Medicine in Residence, University of California at Los Angeles School of Medicine.

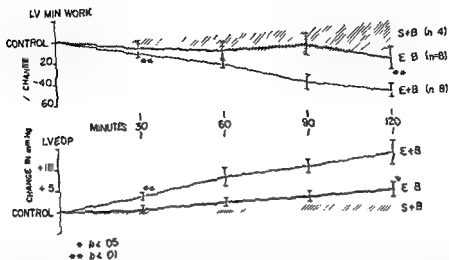


Fig 1 For each group left ventricular minute work (external work = cardiac index \times mean aortic pressure) is expressed as the mean percentage change from control at each period. Left ventricular end diastolic pressure is graphed as the mean change in mm Hg. Limits represent one S.E.M. Significance between the saline animals and each ethanol group was computed with Student's *t* test. S+B = saline with blockade E-B = ethanol without blockade E+B = ethanol with blockade

visual experiments during the first 30 minutes. In Fig 1 the mean changes from control in left ventricular minute work and left ventricular end diastolic pressure are plotted for each group at each period of observation during the infusions. This presentation of the data emphasizes the difference between the two ethanol groups. Without blockade alcohol produced a downward trend in cardiac performance but none of the changes differed from the saline group until 120 minutes. With blockade alcohol produced progressive impairment that was significant by 30 minutes and at all periods greater than in animals without blockade.

Discussion

A comparison of recent canine studies on the hemodynamic consequences of acutely administered ethyl alcohol is complicated by differences in methods which of themselves can influence the overall results profoundly.^{11,12} The studies were conducted in dogs anesthetized with either pentobarbital or alpha-chloralose. Barbiturates directly depress the heart¹³ and both agents have opposite effects on heart rate which are mediated through changes in autonomic balance.¹⁴ All but two studies were carried out in the open-chest preparation

which reduces heart volumes.¹⁵ While this effect is less relevant in preparations with controlled pressure and flow, the thoracotomy itself produces a sympathetic preponderance in autonomic nervous activity.¹⁶ The administration of alcohol also varied from continuous infusions of 15 to 2,000 mg per kilogram of body weight per minute to bolus injections of 0.5 to 5.0 Gm per kilogram of body weight and the blood levels of alcohol achieved when available naturally also varied. Hemodynamic measurements were recorded as early as 5 minutes after the beginning of the alcohol administration and in one series observations continued to 3 hours. Results were expressed either in terms of the Frank-Starling relation of ventricular filling pressure to work or of contractility assessed from strain gauge analysis. The differences in preparation and subsequent dissimilarities in results and interpretation that came from the same laboratory illustrate the importance of methods.^{11,12}

Despite the variation in methods among these studies, a close examination reveals some uniformity of results when the average to concentrated dose schedules were used or where concentrations of blood alcohol rose to at least 1.0 mg per 100 ml. In anesthetized dogs ethanol consistently

Table I Cardiac performance at control and during 2 hour infusions

| | Control | During infusions (minutes) | | | |
|---|-----------|----------------------------|-------------|-------------|-------------|
| | | 30 | 60 | 90 | 120 |
| Cardiac index (ml/min/kg) | | | | | |
| E - B | 117 ± 14 | 111 ± 9 | 102 ± 9† | 101 ± 11† | 93 ± 11‡ |
| L + B | 115 ± 11 | 107 ± 11 | 92 ± 10§ | 76 ± 14§ | 67 ± 12§ |
| S + B | 127 ± 25 | 130 ± 26 | 126 ± 30 | 125 ± 29 | 126 ± 27 |
| Mean aortic pressure (mm Hg) | | | | | |
| E - B | 147 ± 6 | 139 ± 10 | 143 ± 9 | 142 ± 8 | 151 ± 6 |
| E + B | 132 ± 5 | 127 ± 6 | 128 ± 6 | 123 ± 8† | 119 ± 7† |
| S + B | 150 ± 10 | 145 ± 13 | 150 ± 11 | 148 ± 11 | 152 ± 9 |
| Left ventricular end diastolic pressure (mm Hg) | | | | | |
| E - B | 6.1 ± 0.9 | 6.5 ± 1.4 | 7.5 ± 1.3 | 9.3 ± 1.5 | 10.5 ± 1.1* |
| E + B | 7.3 ± 1.3 | 10.8 ± 2.0 | 14.9 ± 3.1* | 17.9 ± 2.5‡ | 20.7 ± 3.8‡ |
| S + B | 9.8 ± 2.1 | 9.5 ± 2.0 | 9.5 ± 2.2 | 10.3 ± 2.1 | 9.8 ± 2.5 |
| Heart rate (beats/min) | | | | | |
| E - B | 106 ± 10 | 105 ± 5 | 95 ± 5 | 97 ± 5 | 102 ± 6 |
| E + B | 139 ± 6 | 136 ± 7 | 139 ± 9 | 135 ± 7 | 134 ± 6 |
| S + B | 133 ± 10 | 132 ± 12 | 132 ± 11 | 135 ± 14 | 138 ± 14 |

Mean ± SEM E - B = ethanol without blockade E + B = ethanol with blockade S + B = saline with blockade

*p < 0.05

†p < 0.01

‡p < 0.005

§p < 0.001

parative reduction in the MAP is most likely fortuitous and is unrelated to the autonomic blockade as the control MAP in the saline with blockade (S+B) dogs was not reduced. Moreover, in 19 animals where the MAP was recorded before and after blockade, the means were essentially the same. The control heart rates for the 2 blocked groups are within one standard deviation of those reported in other blocked dogs anesthetized or awake.^{2,5}

During ethanol infusions cardiac index declined and LVEDP rose in both groups but the changes were more pronounced in the blocked dogs. In the S+B group neither CI nor LVEDP changed significantly throughout the period of observation. Aortic pressure decreased from control only in the E+B group and the change was significant by the 90 minute period. Heart rates remained unchanged from control in all 3 groups.

To assess the effects of autonomic blockade measurements were taken before and after blockade in 19 animals which included all 12 dogs from the 2 blocked groups. Means before and one hour after continuous blockade for cardiac index were 114 and 106 ml per minute per kilogram of body weight (ns) for LVEDP 6.5 and 8.6 mm Hg (p = 0.05) for heart rate 108 and 137 (p < 0.005) and for MAP 130 and 132 mm Hg (ns). Temperatures in the ascending aorta showed no consistent change after blockade and during blockade, varying only 0.5 to 1.0° C in both directions.

While the general trend for both groups of animals receiving ethanol was one of deterioration as judged from a rising LVEDP and a falling cardiac output the changes were more advanced in the E+B group and in the E-B group the changes were both upward and downward in indi-

ments releases catecholamines mainly epinephrine from the adrenal medulla in man and dogs^{28, 29} it would then be available to stimulate the heart through the unblocked beta adrenergic receptors. However the fact that the F-B group does ultimately show significant deterioration suggests that both stimulation of the heart by catecholamines and depression by ethanol were in force during the period of observation. By comparison the E-B group devoid of extrinsic neurohumoral stimuli manifested only the direct depressant effect of ethanol on the heart. Our results are similar to those of Nakano and co-workers¹ who showed that in the open chest preparation the magnitude of myocardial depression from ethanol was markedly greater in dogs treated with reserpine than in untreated dogs.

Thus our findings in the unblocked ethanol group differ but only quantitatively from the results of most of the studies carried out in anesthetized dogs. Although alcohol eventually depressed cardiac performance hemodynamics within the first hour did not change significantly. The apparent unresponsiveness to ethanol was due to individual experiments that showed variations in cardiac output and LVEDP in both directions. Had we chosen a shorter duration of study our conclusions might have been different. These quantitative differences between our results and those of others emphasize the importance of weighing the influence of an inotropic agent on the heart against other determinants of cardiac performance: intrinsic myocardial function, neurohumoral stimuli, the presence of other cardiac stimulants and depressants and the interposition of the investigative methods themselves.

The quantitative difference between our own two ethanol groups leads to the supposition that the negative inotropic response to alcohol can be obscured by extra cardiac stimuli in particular catecholamine release mediated through the autonomic nervous system. This hypothesis can clarify much of the confusion arising from the hemodynamic results reported in human subjects. In these investigations methods and quantities of alcohol used are surprisingly uniform (2 to 6 ounces of 84 to 90 proof) but the subjects are dis-

similar in reference to heart disease.

Four independent laboratories^{21, 24} studied the effects of a single drink of whiskey rum or gin taken by normal subjects at rest and found the cardiac output to be increased over control measurements. Riff and colleagues²¹ and Blomqvist and associates²⁴ further tested their youthful subjects with submaximal and maximal exercise respectively. Both investigators showed that during effort heart rate and cardiac output rose to equivalent levels with and without alcohol.

Catecholamine measurements following alcohol ingestion are sparse but they are uniform in showing an increase.^{30, 31} In humans there is a 67 per cent increase in epinephrine excreted in urine³¹ and in dogs at comparable doses of alcohol per unit of body weight blood epinephrine levels increase two to fivefold.³⁰ Plasma norepinephrine rises only minimally with moderate doses of ethanol. During exercise the increase in plasma catecholamines is from norepinephrine which rises from slightly above to several times over resting concentrations at submaximal to maximal levels of effort.^{3, 36}

According to our supposition the release of epinephrine dominated any negative effect of alcohol in normal subjects at rest. During exercise the generalized discharge of norepinephrine from nerve endings with its potent beta adrenergic stimulation of the heart totally masked any impairment from alcohol and for that matter obscured any weaker beta stimulation from epinephrine release.

By comparison in alcoholics with no clinical evidence of heart disease 6 ounces of 86 proof alcohol produced no changes in heart rate, cardiac output and blood pressure.^{37, 38} Regan and associates³⁷ studied this group further and found that their response to angiotensin infusion was abnormal and when challenged with 12 ounces of alcohol showed a falling stroke output and rising LVEDP.

In subjects with coronary heart disease and stable angina a small dose of alcohol resulted in a significant decline of cardiac output and systolic blood pressure both at rest and during light exercise.³⁹ In another group of subjects with compensated clinical heart disease of different etiologies Gould

caused depression of cardiac performance with one possible exception. In his most recent report Webb and colleagues¹² used a partial bypass preparation and held cardiac output, heart rate and blood pressure constant. Chloralose was the maintenance anesthetic and bilateral vagotomy was employed. At blood levels of alcohol exceeding 600 mg per 100 ml per cent change in dp/dt did not vary and significant changes in LVEDP did not occur, although an illustration showed "more variation in the LVEDP than in most experiments" of approximately ± 10 mm Hg. The conclusion was that ethanol did not have any appreciable effect on myocardial contractility and that the isolated heart-lung preparation minimized peripheral effects of alcohol. In the same preparation and at blood alcohol concentrations of 238 ± 16 mg per 100 ml, Mierzwik and colleagues⁷ found similarly an insignificant decrease in maximum dp/dt associated with a statistically significant increase in LVEDP of only $+2.5$ cm H_2O ; however they concluded that the heart was depressed.

Our methods resembled most closely those of Regan and associates⁴ who used closed chest, anesthetized (pentobarbital) dogs and a continuous infusion of alcohol at a dose less concentrated than ours. By 15 minutes the left ventricle was ejecting a reduced stroke output at a higher LVEDP. In our comparable ethanol with out blockade group (E-B) half of the dogs showed increases in cardiac output and another half showed decreases in LVEDP by 30 minutes. Not until the second hour did the hemodynamics demonstrate significant changes in both cardiac output and LVEDP. These quantitative differences can best be explained by the difference in anesthetic agents used and were observed by Mierzwik and colleagues⁷ who employed chloralose in some dogs and pentobarbital in others.

The earlier and more severe depression of cardiac performance seen in the group with alcohol and blockade must be interpreted in relation to any influence the blocking drugs may have had on the heart. Although large doses of propranolol directly depress myocardial function,^{17,18} smaller beta adrenergic blocking doses have no depressant

effect in hearts already deprived of adrenergic activity,^{3,14,19,20} as is the case in our blocked groups. Atropine, given alone in doses 10 to 5 times smaller than ours, does not depress ventricular function curves in right heart bypass preparations,¹ nor does it change blood pressure, heart rate, or coronary blood flow in denervated hearts.²¹ The direct effect of larger doses of atropine on the myocardium of dogs is unknown. Combined autonomic blockade with both agents in our own series of anesthetized dogs did not alter cardiac output although the heart rate was accelerated. The mean change in LVEDP was $+2.0$ mm Hg and was just significant at the 5 per cent level. Autonomic blockade in the unanesthetized dog with propranolol and vagal cooling changed the heart rate upward and stroke volume downward but not their product; it did not change LVEDP.⁸ Similar results in normal human subjects attest to the lack of depressant effects of pharmacologic isolation of the heart.²²

If autonomic blockade does not reduce cardiac function to what is the difference between our two ethanol groups? Since the directional change for both groups of animals is downward (Fig. 1) the difference between the two represents for the E-B group less depression of cardiac performance in the presence of intact adrenergic and cholinergic functions. While the precise balance between these two functions is unknown at any given moment, the consistent acceleration of heart rate following autonomic blockade in our animals as well as in others²³ suggests that the prevailing nervous activity before blockade in chloralosed dogs and in conscious dogs³ is that of vagal preponderance. Accumulating evidence has corroborated that efferent vagal activity inhibits heart rate and function^{24,25} and conversely that blockade of vagal activity increases ventricular rate and output.²⁷ Thus the presence of intact parasympathetic function in the E-B group does not offer an explanation for its more favorable performance over the E+B group. On the other hand, intact adrenergic function could explain the difference between the two ethanol groups. Since it is known that alcohol in doses both smaller and larger than those used in our experi-

- adrenaline and noradrenaline in the anaesthetized dog *Br J Pharmacol* 26 322 1966
- 15 Ferguson T B Shiple O W and Gregg D W Effect of blood and saline infusion on ventricular end diastolic pressure stroke work stroke volume and cardiac output in the open and closed chest dog *Circ Res* 1 62 1953
- 16 Jose A D and Stitt F Cardiac function after combined beta adrenergic and cholinergic blockade *Circ Res* 21(Suppl III) 231 1967
- 17 Black J W Duncan W A M and Shanks R G Comparison of some properties of propanolol and propranolol *Br J Pharmacol* 25 577 1965
- 18 Shanks R G The pharmacology of beta sympathetic blockade *Am J Cardiol* 18 308 1966
- 19 Flacke J W Osgood P F and Bendixen H H Propranolol and isoproterenol in dogs deprived of sympathetic nerve activity *J Pharmacol Exp Ther* 158:519 1967
- 20 Atanackovic D and Alper M H Beta blocking actions of propranolol in the isolated mammalian heart *Fed Proc* 24 713 1965
- 21 Nayler W G McInnes I Carson V Swann J and Lowe T E The combined effect of atropine and beta adrenergic receptor antagonists on left ventricular function and coronary blood flow *Am Heart J* 77 246 1969
- 22 Essex H E Herrick J F Mann F C and Baldes E J The effect of atropine on the coronary blood flow of trained dogs with denervated and partially denervated hearts *Am J Physiol* 138 683 1943
- 23 Nordenfält I Haemodynamic response to exercise after combined sympathetic and parasympathetic blockade of the heart *Cardiovasc Res* 5 215 1971
- 24 Daggett W M Nugent G C Carr P W Lowers I C and Harada Y Influence of vagal stimulation on ventricular contractility O consumption and coronary flow *Am J Physiol* 212 8 1967
- 25 Degeest H Levy M V Zieske H and Lijman R I Depression of ventricular contractility by stimulation of the vagus nerve *Circ Res* 1 222 1965
- 26 Wildenthal K Mierzwak D S Wyatt H I and Mitchell J H Influence of efferent vagal stimulation on left ventricular function in dogs *Am J Physiol* 216 577 1969
- 27 Stone H L and Bishop V S Ventricular output in conscious dogs following acute vagal blockade *J Appl Physiol* 24:782 1968
- 28 Perman E S The effect of ethyl alcohol on the secretion from the adrenal medulla in man *Acta Physiol Scand* 44 241 1958
- 29 Klingman G I and Goodall McC Urinary epinephrine and levaternol excretion during acute sublethal alcohol intoxication in dogs *J Pharmacol Exp Ther* 121 313 1957
- 30 Siegel J H The effect of enteric ethanol on arterial and portal venous catecholamines *Clin Res* 12:213 1964
- 31 Riff H P Jain A C and Doyle J T Acute hemodynamic effects of ethanol on normal human volunteers *Am Heart J* 78 592 1969
- 32 Blomqvist G Saltin B and Mitchell J H Acute effects of ethanol ingestion on the response to submaximal and maximal exercise in man *Circulation* 42 463 1970
- 33 Juchmes R Hemodynamic effects of ethyl alcohol in man *Am Heart J* 78 133 1969
- 34 Gould L Zahur M Demartino A and Gomprecht R F Cardiac effects of a cocktail *JAMA* 218:1799 1971
- 35 Vendsalu A Studies on adrenaline and noradrenaline in human plasma *Acta Physiol Scand* 49(Suppl 173) 1 1960
- 36 Chodsey C A Harrison D C and Braunwald E Augmentation of plasma norepinephrine response to exercise in patients with congestive heart failure *N Engl J Med* 267 650 1962
- 37 Regan T J Levinson G E Oldewurtel H A Frank M J Weiss A B and Moschos C M Ventricular function in non-cardiacs with alcoholic fatty liver role of ethanol in the production of cardiomyopathy *J Clin Invest* 48:1397 1969
- 38 Wendt V E Ajluni R Bruce T A Prasad A S and Bing R J Acute effects of alcohol on the human myocardium *Am J Cardiol* 17 804 1966
- 39 Conway N Haemodynamic effects of ethyl alcohol in patients with coronary heart disease *Br Heart J* 30 638 1968

and co workers¹⁴ found that only 2 ounces of whiskey diminished cardiac index at rest.

The hemodynamic studies in human subjects can thus be arranged into a continuous spectrum of responses that range from stimulation to depression. The character of the response seems most closely related to the functional capacity of the heart balanced against the presence of both the stimulatory effects of catecholamine release and the depressant effects of ethanol. It is not surprising that different laboratories using low to moderate doses of alcohol have observed (1) an augmentation of cardiac performance in normal subjects, (2) an apparent unresponsiveness in subjects with subclinical heart disease, and (3) a decline of cardiac function in patients with clinical heart disease.

Summary

Experiments from this and other laboratories have indicated that ethanol can both stimulate and depress cardiac function. Surmising that extracardiac stimuli, especially catecholamine release by alcohol, were obscuring its direct effect on the heart, studies were combined with autonomic blockade in anesthetized closed chest dogs. Ethanol was infused at rates of 25 to 33 mg per kilogram of body weight per minute. Autonomic blockade was achieved with an initial dose of atropine 1.0 mg per kilogram of body weight and propranolol 0.2 mg per kilogram of body weight and maintained by constant infusion. Cardiac performance was assessed from the Frank-Starling relation of ventricular filling pressure to cardiac work at control and at intervals after the start of infusions. Saline infusion with blockade caused no change in cardiac performance over 120 minutes. Ethanol with blockade caused progressive falls from 30 minutes on. Alcohol without blockade produced rises and falls in different animals in the first 30 minutes; thereafter cardiac performance declined but at all intervals less than in dogs given alcohol with blockade.

We concluded that alcohol uniformly depressed the heart after autonomic blockade but without blockade this effect could be masked by extracardiac mechanisms mediated through the autonomic system. Assuming that catecholamine release was

an important extracardiac mechanism, the conclusion was tested against conflicting reports in the literature and was found to be a useful hypothesis.

I acknowledge with thanks technical assistance from Morris Berger, Robert Cullison, Albert Skulsky, and Sam Snyder, and secretarial help from Kwei Chou. I am indebted to the late Anthony D. Jose, M.D., for his thought-provoking discussions.

REFERENCES

1. Mitchell J. H. and Cohen I. S. Alcohol and the heart. *Mod. Concepts Cardiovasc. Dis.* 39:109, 1970.
2. Jose A. D. and Stitt F. Effects of hypoxia and metabolic inhibitors on the intrinsic heart rate and myocardial contractility in dogs. *Circ. Res.* 25:53, 1969.
3. Jose A. D. and Taylor R. R. Autonomic blockade by propranolol and atropine to study intrinsic myocardial function in man. *J. Clin. Invest.* 48:2019, 1969.
4. Regan T. J., Koroxenidis G., Moschos C. B., Oldewurtel H. A., Lehn P. H., and Hellem H. K. The acute metabolic and hemodynamic responses of the left ventricle to ethanol. *J. Clin. Invest.* 45:270, 1966.
5. Bishop V. S. and Horwitz I. D. Effects of altered autonomic control on left ventricular function in conscious dogs. *Am. J. Physiol.* 22:1278, 1971.
6. Ganz V. The acute effect of alcohol on the circulation and on the oxygen metabolism of the heart. *Am. Heart J.* 66:494, 1963.
7. Mierzwik D. S., Wildenthal K., and Mitchell J. H. Effect of ethanol on the canine left ventricle. *Clin. Res.* 15:215, 1967.
8. Nakano J., Kessinger J. M., and Prince A. V. Cardiovascular effects of ethanol and the adrenergic mechanism. *Clin. Res.* 19:24, 1971.
9. Newman W. H. and Valicenti J. F., Jr. Ventricular function following acute alcohol administration. A strain gauge study of depressed ventricular dynamics. *Am. Heart J.* 81:61, 1971.
10. Webb W. R. and Degerli I. U. Ethyl alcohol and the cardiovascular system. *J.AMA* 191:1055, 1965.
11. Webb W. R., Degerli I. U., Cook W. A., and Ural M. D. Alcohol, digitalis, and cortisol and myocardial capacity in dogs. *Ann. Surg.* 163:811, 1966.
12. Webb W. R., Gupta D. N., Cook W. A., Sugg W. I., Brishour F. A., and Ural M. O. Effects of alcohol on myocardial contractility. *Dis. Chest* 52:602, 1967.
13. Daniel F. E., Fulton J. B., Hiddleston M., Martin W., and Foulks J. G. Analysis of the mechanism of barbiturate induced cardiovascular depression and its antagonism by sympathomimetic amines. *Arch. Intern. Pharmacodyn.* 108:457, 1956.
14. Shanks R. G. The effects of propranolol on the cardiovascular responses to isoprenaline.

Investigation of atrial aberration as a cause of altered P wave contour

Peter Probst MD

Jane Hunter, AB

Olive Gamble AB

Keith Cohn, MD

San Francisco, Calif

Explaining a changing P wave contour is oftentimes the bane of the electrocardiographer. Sudden variation in P wave morphology is generally due to a change in the site of atrial focus such as occurs in *atrial premature contractions* or *atrial escape beats*. Gradual variations in P wave contour are alternatively explained on the basis of a *wandering pacemaker* whereby it is assumed that the focus of atrial excitation moves gradually from one portion of the atrium to another. *Atrial fusion* is characterized by depolarization of the atrium via two origins simultaneously—usually the sinus and an ectopic focus—and this is yet another mechanism of changing P wave morphology. One additional mechanism proposed to explain sudden changes in P wave contour—and that which is the subject of this study—is aberrant impulse conduction through the atrium (*atrial aberration*).^{1,2} Here, it is proposed that the site of impulse formation remains fixed but because portions of the atria are partially or totally refractory when the wave front reaches them, the impulse is forced to take an alternate course resulting in morpho-

logical changes in the P wave contour. This particular mechanism may be considered analogous to aberrant ventricular conduction whereby gross changes in the QRS appearance come about when refractory fascicular, Purkinje or ventricular tissue is encountered by the depolarizing wave front.^{3,4} By infringing on the refractory period of the previous beat, then premature supraventricular contractions or runs of supraventricular tachycardia could be aberrantly conducted through the atrium or ventricle resulting in variation of P wave morphology or in a distorted QRS complex. Aberration of an escape beat—one which follows a pause—has also been described.⁵

It should be pointed out that morphological changes in P wave contour also occur consequent to pathological changes within the atrium or atrial enlargement but such distorted P waves tend to be 'persistent'—present in all the beats—rather than being intermittent.

Although ventricular aberration is an unequivocal cause of striking changes in QRS duration and morphology variations

From The Department of Cardiology, Presbyterian Hospital, Pacific Medical Center and The Heart Research Institute, The Institute of Medical Sciences, Pacific Medical Center, San Francisco, Calif.
This study was supported by Grant HL 05498 from the National Institutes of Health.
This study was published Dec 19, 1972.
Reprint requests to Keith Cohn, MD, Cardiopulmonary Laboratories, Pacific Medical Center, P.O. Box 7999, San Francisco, Calif 94120.

in P wave contour resulting from this mechanism are more speculative and the phenomenon of atrial aberration ■ yet to be investigated systematically This study then was undertaken in order to test whether aberrant conduction within the atrium ■ in fact capable of altering P wave morphology and if such is the case whether the changes in P wave contour are marked—producing a total reversal of polarity—or only slight resulting in minor variations in P wave duration or appearance It is presumed that information derived from this study will enhance an understanding of atrial electrophysiology and should aid the clinician in comprehending complex electrocardiographic tracings

Method

Twenty three healthy adult mongrel dogs weighing 15 to 23 kilograms were anesthetized with sodium pentobarbital (30 mg per kilogram of body weight intravenously) Respiration was maintained through a cuffed endotracheal tube attached to an automatic volume respirator pump One femoral vein was cannulated for the injection of drugs a pressure catheter was inserted into the right atrium via the other femoral vein and a femoral artery was cannulated for monitoring blood gases and pH Right or bilateral thoracotomies in the fourth intercostal space were performed and the heart was suspended in a pericardial sling Electrodes composed of gold tipped stainless steel needles fixed in plastic 2 mm apart were sutured with 4 0 cardiovascular silk to five sites on the epicardial surface of the right atrium (1) the region of the sino atrial node (2) the junction of the coronary sinus with the right atrium (3) the junction of the superior vena cava with the right atrium (4) the junction of the inferior vena cava with the right atrium and (5) the interior mid portion of the VV groove These multiple electrodes were employed to observe lateral longitudinal and anteroposterior direction of the vector in order to enhance the possibilities of detecting any change in P wave morphology The electrodes at the sinoatrial and coronary sinus sites acted as bipolar pacing electrodes and those from the vena cavae and AV groove served as

unipolar recording electrodes In three dogs additional electrodes were secured on the left atrial surface at three sites pacing and recording electrodes at the center of the free wall of the left atrium and another recording electrode at the left atrial portion of Bachmann's bundle Standard limb Leads I II and III were also monitored

S₁ and S₂ Grass pulse generators connected through Grass isolation units and constant current units delivered both the driving and premature pulses through the same electrode A switch that simultaneously turned off the driving stimulus (S₀) triggered the atrial premature stimuli either singly (S₁) or in pairs (S₁ and S₂) The S₁ impulse was positioned at the S₀ effective refractory period (ERP) by moving the S₁ stimulus in at 5 msec intervals until it no longer produced a propagated response S₂ was positioned at the S₁ ERP in the same manner The stimulus strength for S₀ S₁ and S₂ was set at 2 Ma above threshold threshold was defined as the minimum amperage necessary to drive the heart at a given rate Only paced premature and postextrasystolic beats were evaluated for aberration to eliminate atrial ectopy as causes of P contour change Atrial fusion was also avoided by having the paced beats precede unpaced ones

Because the premature P often hit upon the T wave of the preceding beat change in its configuration was difficult to evaluate For this reason complete heart block was produced in 10 dogs by making an incision through the AV node The production of complete heart block allowed the delivery of P waves late in ventricular diastole after completion of the T wave A purse string suture of 3 0 silk was secured around the right atrial appendage and the tip of the appendage was removed to provide entrance into the right atrium A small knife designed to fit flat against the index finger was inserted the site of the AV node located by palpation and a cut approximately 1 cm long was made through the AV node More than one cut was made if the first failed to produce complete heart block The hearts were examined after the study was terminated and in seven hearts the cut through the AV node extended to the atrial septum Comparing these studies with the three in which the

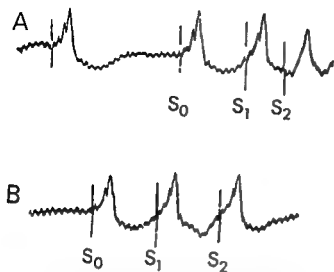


Fig 1 *A* and *B* Electrically stimulated P waves are recorded from Lead II. *A* the P wave of the first premature stimulus (S_1) does not differ significantly in configuration from the normally paced beats (S_0). The P wave following the second premature stimulus (S_2) is slightly smaller and less notched. *B* Lesser degrees of prematurity result in the abolition of atrial aberration. Complete heart block is present in both tracings and QRS complexes are not seen.

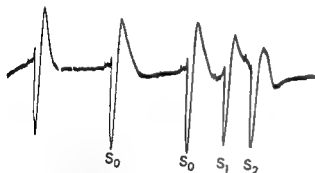


Fig 2 Atrial electrograms taken when the region of the sinoatrial node is stimulated; the recording is being taken from the region of Bachmann's bundle. The P waves following both the first and second premature stimuli (S_1 , S_2) differ slightly in contour from the normally paced beats (S_0) and also differ slightly from each other. Complete heart block is present and QRS complexes are not evident. Note the atrial repolarization waves.

septum was not damaged yielded no significant differences in the frequency or degree of aberration, so the 10 studies are considered together. After AV block was obtained a pulse of 2 msec duration drove the atria at a rate of three to four cycles per second; the rate and duration were held constant throughout the experiment. The right ventricle was intermittently paced at a rate of 1.5 to 2.5 cycles per second to maintain the heart's viability.

In four dogs, additional studies in the normally conducting heart were undertaken to clarify the role of right atrial pressure (or myocardial stretch) on P wave contour. Two methods were used to increase atrial pressure: (1) rapidly injecting four consecutive 100 ml boluses of Ringer's lactate solution into the right atrium through the femoral venous catheter, raising the pressure from between 6 and 14 mm Hg, and (2) inserting a balloon through the wall of the right ventricle and gradually inflating it. Paced and un-paced recordings were taken during the injection and balloon inflation to evaluate the effect of these maneuvers on P wave configuration when the cycle length was held constant.

All recordings were made on a Honeywell multichannel photographic recorder. Two channels recording atrial and one standard limb lead were recorded simultaneously.

Changes in P wave morphology were described in terms of amplitude, contour, and duration. The effects of five variables on atrial aberration were considered: (1) the degree of prematurity of the premature beats, evaluated in all studies by comparing the premature beats (S_1 and S_2) to each other and to the paced beat (S_0); (2) stimulus strength, considered in five studies by comparing S_1 and S_2 at the standard stimulus and at four to tenfold multiples of the standard strength; (3) influence of differing stimulus sites; (4) influence of changing heart rate (atrial pacing rate); and (5) effect of increasing atrial pressure or stretch on P wave configuration.

Results

Complete heart block preparations. The P wave contour could be studied most precisely—without becoming superimposed on the preceding T wave—under conditions of complete heart block. In these studies only minor changes in P wave amplitude, duration, and contour appeared as the prematurity became greater and the premature P wave was made to impinge upon the effective refractory period (ERP) of the preceding beat. In limb Leads I, II, and III we observed amplitude changes ranging from -0.093 to $+0.15$ mv (± 9.9 per cent), duration changes ranged from -10 to $+12$ msec (± 6 per cent), and slight

contour changes such as minor degrees of notching were elicited in the premature impulse (S_1) (Figs 1 and 2). As the S_0 S_1 coupling interval was gradually widened the changes in amplitude, duration and contour progressively became less marked until the premature beat appeared to be identical with the paced beat (Fig 1). It should be emphasized that the changes in P wave contour noted were slight, often absent and there were no instances where P wave polarity became reversed—i.e. when a negative P wave became positive or vice versa.

Introducing pairs of premature stimuli (S_1 S_2) also elicited minor changes in amplitude (-0.26 to $+0.22$ mv, ± 14 per cent change), duration (-20 to $+10$ msec) and minor degrees of notching seen in the limb leads. Although the configurations of the first and second premature P waves often differed slightly from one another, the overall changes continued to be slight and still no marked variation in P wave contour or reversal in polarity was detected (Figs 1 and 2). When the atrial pacing rate was suddenly increased the initial P waves showed minor changes similar to those described above; however, the subsequent beats reverted to the original P wave morphology.

Increasing the strength of the premature impulse by 4 to 10 times stimulus threshold permitted a greater degree of prematurity since further penetration into the S_0 effective refractory period could be accomplished and this resulted in amplitude, duration and contour changes slightly exceeding those described above. However, variations in the appearance of the P wave remained notably slight. Increasing the strength of the premature stimulus without changing coupling interval did not affect P wave morphology.

It was thought possible that certain pathways within the atrium might be more vulnerable to the development of atrial aberration than others. This hypothesis was looked into by stimulating the atrium from three different electrodes. No particular pattern was evident in these studies—i.e. the frequency and minor degrees of aberration seen were the same whichever stimulus sites were employed.

Finally, it was noted that slight changes

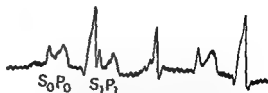


Fig 3 Lead II recorded in a preparation which has no complete heart block. The premature P wave (P) is shown to manifest slight aberration, being smaller and slightly more peaked. The larger complexes are QRS's.

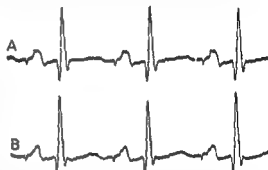


Fig 4 The P wave contour is seen before (A) and after (B) inflating a balloon in the right ventricle. Note that the P wave becomes slightly more peaked and the notch moves onto its upstroke.

in the pacing (S_0) P wave configuration occurred with the production of complete heart block. In 7 of the 10 studies the induction of complete block resulted in slight injury to the atrial septum. The results from these studies were indistinguishable from those where the septum remained intact, leaving us to conclude that septal injury per se did not influence the degree of aberration observed.

Absence of complete heart block: In the studies in the normally conducting heart we evaluated both the premature and postextrasystolic beat for aberration. Again, both types of beats showed only slight—at times negligible—changes in amplitude, contour and duration (Fig 3). Oftentimes the beats fell upon the T wave of the preceding beat, making it difficult to determine whether prematurity or superimposition on the T wave was more responsible for the contour change. In those studies in which the T wave was small or isoelectric, the I change could be quantitated as decreases in P wave duration up to 5 msec and/or amplitude up to 0.10 mv.

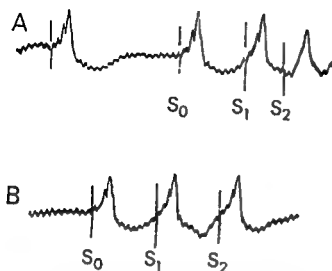


Fig 1 *A* and *B* Electrically stimulated P waves are recorded from Lead II. In the P wave of the first premature stimulus (S_1) does not differ significantly in configuration from the normally paced beats (S_0). The P wave following the second premature stimulus (S_2) is slightly smaller and less notched. *B* Lesser degrees of prematurity result in the abolition of atrial aberration. Complete heart block is present in both tracings and QRS complexes are not seen.

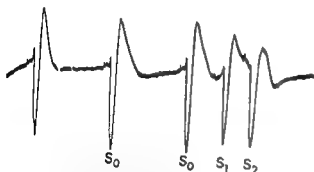


Fig 2 Atrial electrograms taken when the region of the sinoatrial node is stimulated; the recording is being taken from the region of Bachmann's bundle. The P waves following both the first and second premature stimuli (S_1 , S_2) differ slightly in contour from the normally paced beats (S_0) and also differ slightly from each other. Complete heart block is present and QRS complexes are not evident. Note the atrial repolarization waves.

septum was not damaged yielded no significant differences in the frequency or of degree of aberration, so the 10 studies are considered together. After AV block was obtained, a pulse of 2 msec duration drove the atria at a rate of three to four cycles per second; the rate and duration were held constant throughout the experiment. The right ventricle was intermittently paced at a rate of 1.5 to 2.5 cycles per second to maintain the heart's viability.

In four dogs additional studies in the normally conducting heart were undertaken to clarify the role of right atrial pressure (or myocardial stretch) on P wave contour. Two methods were used to increase atrial pressure: (1) rapidly injecting four consecutive 100 ml boluses of Ringer's lactate solution into the right atrium through the femoral venous catheter raising the pressure from between 6 and 14 mm Hg, and (2) inserting a balloon through the wall of the right ventricle and gradually inflating it. Paced and un-paced recordings were taken during the injection and balloon inflation to evaluate the effect of these maneuvers on P wave configuration when the cycle length was held constant.

All recordings were made on a Honeywell multichannel photographic recorder. Two channels recording atrial and one standard limb lead were recorded simultaneously.

Changes in P wave morphology were described in terms of amplitude, contour, and duration. The effects of five variables on atrial aberration were considered: (1) the degree of prematurity of the premature beats, evaluated in all studies by comparing the premature beats (S_1 and S_2) to each other and to the paced beat (S_0); (2) stimulus strength considered in five studies by comparing S_1 and S_2 at the standard stimulus and at four to tenfold multiples of the standard strength; (3) influence of differing stimulus sites; (4) influence of changing heart rate (atrial pacing rate); and (5) effect of increasing atrial pressure or stretch on P wave configuration.

Results

Complete heart block preparations The P wave contour could be studied most precisely—without becoming superimposed on the preceding T wave—under conditions of complete heart block. In these studies only minor changes in P wave amplitude, duration, and contour appeared as the prematurity became greater and the premature P wave was made to impinge upon the effective refractory period (ERP) of the preceding beat. In limb Leads I, II, and III we observed amplitude changes ranging from -0.093 to $+0.15$ mv (± 9.9 per cent), duration changes ranged from -0.10 to $+12$ msec (± 6 per cent), and slight

roversy as to whether impulse propagation through the atrium depends upon the specialized interatrial tracts or whether atrial depolarization simply progresses in a quasi radial fashion corresponding to the gross anatomical landmarks.¹¹ We reasoned that assuming these tracts are physiologically significant they would in all likelihood have nonidentical refractory periods. Thus a sufficiently premature stimulus might encounter a pathway which was incapable of normal conduction and the morphology of the P wave would thereby be grossly altered. This is precisely what is seen in ventricular aberration where gradual increments in prematurity often lead to the sudden appearance of right bundle branch block.* We therefore compared the effects of different sites of stimulation on the degree of atrial aberration speculating that an impulse originating from a particular locus might manifest a more marked degree of aberration were it to normally utilize an atrial pathway and were it to encounter refractoriness of that particular tract.¹² The results of this portion of the study were clear only *gradual* or *slight changes* in P wave morphology were detected in our study as the degree of prematurity was increased and the magnitude of change was uninfluenced by location of the stimulating impulse. These findings therefore indirectly suggest that the internodal tracts play only a minor role in impulse spread through the atria. An alternative explanation of these data would be that the refractory period of all the tracts and of the atrial musculature are essentially uniform. Thus the evidence that these tracts are physiologically important (P wave alterations are seen in experimental animals after focal lesions have been produced at sites of interatrial tracts¹³) cannot be negated by these findings.

The second observation of note was that increased atrial fiber stretch or pressure—in the presence of a fixed heart rate—produced essentially the same morphological

P wave alterations as did varying cycle length. A premature atrial contraction often occurs when the AV valves are closed or when the ventricular relaxation is incomplete suggesting that this mechanical effect on P wave contour might be as important as the temporal effect. Transient alterations in P wave configuration also occur in certain clinical situations such as acute cor pulmonale status asthmaticus and in left ventricular failure following acute myocardial infarction—conditions where an atrium is temporarily dilated. The mechanism by which atrial stretch or pressure influences the appearance of the P wave is unknown and might include slight rotations of the heart or the effect of atrial stretch on cellular membrane potential or atrial conduction. Whatever the underlying mechanism one is forced to conclude that the aberrant atrial beats detected in this study might be related either to their timing or to varying fiber tension or stretch.

The final point of emphasis is that the changes we observed in P wave morphology were slight much less in degree than most of the P wave alterations we usually try to explain in clinical electrocardiography. Thus were we to extrapolate from animal studies to clinical situations—a step that should be taken with some caution—we would conclude that most of the gross changes in P wave contour are related to factors other than aberrancy: wandering atrial pacemaker, atrial or junctional ectopy, fusion beats, respiratory effects or artifact. Clinical electrocardiography remains sufficiently speculative so that it would seem unwise at this time to invoke a mechanism such as atrial aberration to explain all but the most minor alterations in P wave configurations seen.

Summary

In order to study atrial aberration atrial premature beats and postextrasystolic beats were electrically induced in 23 dogs and the P wave contour was recorded in limb and atrial leads. In these studies the site and timing of stimulation were controlled to eliminate ectopy and fusion as causes of P wave change. Further P wave analysis was facilitated by producing complete AV block in 10 dogs thereby

appeared in the premature or postextrasystolic beat

Right atrial volume and pressure change
When right atrial volume was suddenly or gradually increased by volume expansion or by inflating a balloon in the right ventricle alterations in P wave duration, amplitude, and contour similar to those seen during premature stimuli were noted, even though cycle length was held constant (Fig 4). Duration varied by ≈ 10 msec, amplitude changed by ≈ 14 mv, and again only slight degree of P wave notching without reversal of polarity was detected. The greatest changes in P wave morphology were produced with the greater degrees of pressure and volume

Discussion

This study was designed to determine whether aberrant conduction through the atrium might be one mechanism of producing transient alterations in P wave morphology. Before delving into the subject further it is important to establish certain definitions. Aberrancy is the term generally applied to describe temporary alterations in P or QRS configuration brought about when an impulse reaches tissue which is refractory and therefore incapable of allowing normal conduction. Excluded from the definition of aberrancy then are alterations in P or QRS configuration due to ectopic foci, to fusion, or to more permanent conduction disturbances consequent to myocardial degeneration, infiltration, or fibrosis.

Ventricular aberration is a well accepted entity and occurs when a supraventricular impulse encounters an incompletely repolarized bundle branch or fascicle.^{2,5} This most commonly is seen in a premature beat or during tachycardia where the shortened cycle length enhances the likelihood that the impulse will infringe upon the refractory period of the preceding beat. Cardiac slowing also increases the possibility that a subsequent premature beat (with a shortened cycle length) will encounter refractory fibers, so that the combination of a long interval followed by a short interval constitutes the most favorable set of circumstances for the production of ventricular aberration.^{2,5} Since the refractory period of the right bundle branch

exceeds that of the AV node, His bundle and left bundle branch, functional right bundle branch block is the most common form of ventricular aberration seen. Paradoxical aberration, the appearance of a distorted QRS complex which appears following a pause is observed less commonly but remains a well established entity.⁵

The concept of aberrant atrial conduction has been invoked to explain bizarre configurations of the sinus P wave following premature atrial, AV junctional and ventricular contractions.^{1,2} Chung and colleagues' studies^{1,2} consisted of reviewing clinical electrocardiograms and the P waves considered to show atrial aberration were those terminating the postextrasystolic pause. We presume that these P waves, rather than the premature ones were studied since in clinical electrocardiography one is unable to exclude atrial ectopy as a cause of P wave alteration in premature beats. Thus although atrial aberration is more likely to occur during a shortened cycle length, Chung and colleagues^{1,2} were obliged to concentrate only on the P waves ending a prolonged cycle in their search for atrial aberration. One should point out at this juncture that even under these circumstances ectopy or fusion of the escaped beat cannot be excluded as a potential cause of altered postextrasystolic P wave configuration.

The conditions were set in our present study to optimize the chances of detecting changes in P wave configuration due to aberrant conduction. At the same time controlling factors were built in to eliminate atrial ectopy, wandering pacemaker and atrial fusion as potential complicating factors which might influence P wave contour. Under these controlled conditions only minor changes in P wave morphology and duration were observed in marked abnormalities such as reversal of P wave polarity were never seen. Thus although cycle length was varied and excessive prematurity produced (to the point of reaching complete atrial refractoriness) and beats following postextrasystolic pauses were studied only minimal degrees of aberrant atrial conduction were seen.

This study touches on two other points of interest. The first concerns the current con-

troversty as to whether impulse propagation through the atrium depends upon the specialized interatrial tracts or whether atrial depolarization simply progresses in a quasi radial fashion corresponding to the gross anatomical landmarks.^{8,9} We reasoned that assuming these tracts are physiologically significant they would in all likelihood have nonidentical refractory periods. Thus a sufficiently premature stimulus might encounter a pathway which was incapable of normal conduction and the morphology of the P wave would thereby be grossly altered. This is precisely what is seen in ventricular aberration where gradual increments in prematurity often lead to the sudden appearance of right bundle branch block.⁸ We therefore compared the effects of different sites of stimulation on the degree of atrial aberration speculating that an impulse originating from a particular locus might manifest a more marked degree of aberration were it to normally utilize in atrial pathway and were it to encounter refractoriness of that particular tract.¹⁰ The results of this portion of the study were clear: only *gradual* or *slight* changes in I wave morphology were detected in our study as the degree of prematurity was increased and the magnitude of change was uninfluenced by location of the stimulating impulse. These findings therefore indirectly suggest that the internodal tracts play only a minor role in impulse spread through the atria. An alternative explanation of these data would be that the refractory period of all the tracts and of the atrial musculature are essentially uniform. Thus the evidence that these tracts are physiologically important (P wave alterations are seen in experimental animals after focal lesions have been produced at sites of interatrial tracts¹¹) cannot be negated by these findings.

The second observation of note was that increased atrial fiber stretch or pressure—in the presence of a fixed heart rate—produced essentially the same morphological

P wave alterations as did varying cycle length. A premature atrial contraction often occurs when the AV valves are closed or when the ventricular relaxation is incomplete suggesting that this mechanical effect on P wave contour might be as important as the temporal effect. Transient alterations in P wave configuration also occur in certain clinical situations such as acute cor pulmonale status asthmaticus and in left ventricular failure following acute myocardial infarction—conditions where an atrium is temporarily dilated. The mechanism by which atrial stretch or pressure influences the appearance of the P wave is unknown and might include slight rotations of the heart or the effect of atrial stretch on cellular membrane potential or atrial conduction. Whatever the underlying mechanism, one is forced to conclude that the aberrant atrial beats detected in this study might be related either to their timing or to varying fiber tension or stretch.

The final point of emphasis is that the changes we observed in P wave morphology were slight, much less in degree than most of the P wave alterations we usually try to explain in clinical electrocardiography. Thus we were to extrapolate from animal studies to clinical situations—a step that should be taken with some caution—we would conclude that most of the gross changes in P wave contour are related to factors other than aberrancy: wandering atrial pacemaker, atrial or junctional ectopy, fusion beats, respiratory effects or artifact. Clinical electrocardiography remains sufficiently speculative so that it would seem unwise at this time to invoke a mechanism such as atrial aberration to explain all but the most minor alterations in P wave configurations seen.

Summary

In order to study atrial aberration, atrial premature beats and postextrasystolic beats were electrically induced in 23 dogs and the P wave contour was recorded in limb and atrial leads. In these studies the site and timing of stimulation were controlled to eliminate ectopy and fusion as causes of P wave change. Further P wave analysis was facilitated by producing complete AV block in 10 dogs thereby

Admittedly, the results of this study are limited by the fact that the degree of prematurity was not varied systematically. The results of this study were clear: only gradual or slight changes in I wave morphology were detected in our study as the degree of prematurity was increased and the magnitude of change was uninfluenced by location of the stimulating impulse. These findings therefore indirectly suggest that the internodal tracts play only a minor role in impulse spread through the atria. An alternative explanation of these data would be that the refractory period of all the tracts and of the atrial musculature are essentially uniform. Thus the evidence that these tracts are physiologically important (P wave alterations are seen in experimental animals after focal lesions have been produced at sites of interatrial tracts¹¹) cannot be negated by these findings.

appeared in the premature or postextra systolic beat

Right atrial volume and pressure change
When right atrial volume was suddenly or gradually increased by volume expansion or by inflating a balloon in the right ventricle, alterations in P wave duration, amplitude and contour similar to those seen during premature stimuli were noted even though cycle length was held constant (Fig 4). Duration varied by ± 10 msec, amplitude changed by ± 14 mv and again only slight degree of P wave notching without reversal of polarity was detected. The greatest changes in P wave morphology were produced with the greater degrees of pressure and volume.

Discussion

This study was designed to determine whether aberrant conduction through the atrium might be one mechanism of producing transient alterations in P wave morphology. Before delving into the subject further it is important to establish certain definitions. Aberrancy is the term generally applied to describe temporary alterations in P or QRS configuration brought about when an impulse reaches tissue which is refractory and therefore incapable of allowing normal conduction. Excluded from the definition of aberrancy then, are alterations in P or QRS configuration due to ectopic foci to fusion or to more permanent conduction disturbances consequent to myocardial degeneration infiltration or fibrosis.

Ventricular aberration is a well accepted entity and occurs when a supraventricular impulse encounters an incompletely repolarized bundle branch or fascicle.^{2,3} This most commonly is seen in a premature beat or during tachycardia where the shortened cycle length enhances the likelihood that the impulse will infringe upon the refractory period of the preceding beat. Cardiac slowing also increases the possibility that a subsequent premature beat (with a shortened cycle length) will encounter refractory fibers so that the combination of a long interval followed by a short interval constitutes the most favorable set of circumstances for the production of ventricular aberration.^{2,3} Since the refractory period of the right bundle branch

exceeds that of the AV node, His bundle, and left bundle branch, functional right bundle branch block is the most common form of ventricular aberration seen. *Paradoxical aberration* the appearance of a distorted QRS complex which appears following a pause is observed less commonly but remains a well established entity.⁵

The concept of *aberrant atrial conduction* has been invoked to explain bizarre configurations of the "sinus" P wave following premature atrial AV junctional, and ventricular contractions.^{1,2} Chung and colleagues' studies^{1,2} consisted of reviewing clinical electrocardiograms, and the P waves considered to show atrial aberration were those terminating the postextra systolic pause. We presume that these P waves rather than the premature ones were studied since in clinical electrocardiography one is unable to exclude atrial ectopy as a cause of P wave alteration in premature beats. Thus although atrial aberration is more likely to occur during a shortened cycle length, Chung and colleagues^{1,2} were obliged to concentrate only on the P waves ending a prolonged cycle in their search for atrial aberration. One should point out at this juncture that even under these circumstances ectopy or fusion of the escaped beat cannot be excluded as a potential cause of altered postextrasystolic P wave configuration.

The conditions were set in our present study to optimize the chances of detecting changes in P wave configuration due to aberrant conduction. At the same time controlling factors were built in to eliminate atrial ectopy wandering pacemaker, and atrial fusion as potential complicating factors which might influence P wave contour. Under these controlled conditions only minor changes in P wave morphology and duration were observed marked abnormalities such as reversal of P wave polarity were never seen. Thus although cycle length was varied and excessive prematurity produced (to the point of reaching complete atrial refractoriness) and beats following postextrasystolic pauses were studied only minimal degrees of aberrant atrial conduction were seen.

This study touches on two other points of interest. The first concerns the current con-

Viral infection of the aorta of man associated with early atherosclerotic changes

G E Burch MD
J M Harb PhD
Y Hiramoto MD
Lana Shewey MD
New Orleans La

The production of experimental atherosclerosis has been accomplished in numerous primates and other species by inducing hypertension and by dietary production of hyperlipemia.¹ Furthermore attempts have been made to study the regression of aortic fatty lesions following cessation of high cholesterol diets.^{2,4} The most prominent feature of such atherosclerotic lesions whether progressive or regressive has been the presence of lipid-containing cells in the aortic intima and in the inner elastic layers of the media. The lipid-containing cells were considered by some investigators to be monocytes which had infiltrated into the aortic intima,⁵ whereas others have considered that foam cells are the predominant lipid-containing cells in experimentally induced atherosclerotic lesions.⁶ In proliferative (non necrotic) lesions in monkeys the predominant cells observed were smooth muscle cells with fewer cells resembling fibroblasts.⁶

The purpose of this report is to describe the finding of lipid-containing cells in the

intima muscularis and internal elastic layer of the media of the aorta of a patient who had viral cardiomyopathy most probably due to the Coxsackie B₁ virus. Lipid deposition suggestive of atherosclerotic alteration was found in the aorta of this patient. It is suggested that the lipid deposition occurred as a response to a viral infection of the aorta itself an initiating stimulus other than hypertension or high cholesterol diet.

Case report

This 19 year old man was in excellent health until two weeks prior to his admission to hospital in Monroe Louisiana, on February 23 1972 for congestive heart failure. He had gradually developed swelling of the abdomen and feet with mild dyspnea following an episode of diarrhea nausea and vomiting after eating pork on February 9 1972. Other than a past history of familial asthma treated with Primatene most of his life there was no apparently significant illness. However he had had the usual respiratory diseases. Examination on admission to hospital revealed generalized cardiomegaly loud first and second mitral valve sounds protodiastolic gallop rhythm and a soft systolic blowing murmur along the lower left sternal border. There was a trace of edema of the ankles with elevated venous pressure

From the Department of Medicine of the Tulane University School of Medicine and the Charity Hospital of Louisiana, New Orleans, La.
Supported by Grant HL-06769 from the National Heart and Lung Institute of the United States Public Health Service and the Rudolph M. Isaacs Memorial Fund for the Late President Hiram Laboratry the Rowell A. Billups Fund for Research in Heart Disease and the F. and L. Labatry.
Received for publication February 27 1973.
Reprint requests to Dr. George E. Burch, 1430 Tulane Avenue, New Orleans, La. 70112.

preventing superimposition of the premature P on the preceding T wave

As the coupling interval was progressively narrowed, minor changes in the premature P duration, amplitude, and contour (such as slight degrees of notching) appeared. The postextrasystolic P wave manifested even lesser changes. Varying the myocardial stretch by changing right atrial volume produced similar alterations in P wave configuration at a constant cycle length. Comparable variations in atrial stretch occur with premature beats, and these mechanical effects may, therefore, be one mechanism of the P contour change.

Atrial aberration was uninfluenced by site or strength of stimulus and was minimally affected by rate. No reversal in P wave polarity was observed.

We conclude that only small alterations in P configuration result from atrial aberration and that this mechanism should not be invoked to explain any but the most minor alterations in P wave configuration.

REFERENCES

- 1 Chung E K. Aberrant atrial conduction: Unrecognized electrocardiographic entity. *Br Heart J* 34:341, 1972.
- 2 Chung E K, Walsh T J, and Mason F. Atrial parasystole. *Am J Cardiol* 14:255, 1964.
- 3 Katz I N and Pick A. Clinical electrocardiography, part I. The arrhythmias. Philadelphia 1956. Lea & Febiger Publishers.
- 4 Marriott H J L and Sandler I A. Criteria old and new for differentiating between ectopic ventricular beats and aberrant ventricular conduction in the presence of atrial fibrillation. *Progr Cardiovasc Dis* 9:18, 1966.
- 5 Singer D H and Ten Eick R E. Aberrancy: electrophysiologic aspects. *Am J Cardiol* 28:381, 1971.
- 6 James T N. The connecting pathways between the sinus node and A V node and between the right and the left atrium in the human heart. *Am Heart J* 66:498, 1963.
- 7 Durrer D, Van Dam R Th, Freud G E, Janse M J, Meijler F L, and Arzbaeher R C. Total excitation of the isolated human heart. *Circulation* 41:899, 1970.
- 8 Spach M S, Lieberman M, Scott J G, Barr R C, Johnson E A, and Kootsey J M. Excitation sequences of the atrial septum and the A V node in isolated hearts of the dog and rabbit. *Circ Res* 29:156, 1971.
- 9 Goodman D, Van Der Steen A B M, and Van Dam R Th. Endocardial and epicardial activation pathways of the canine right atrium. *Am J Physiol* 220:1, 1971.
- 10 Waldo A L, Vitikainen K J, Kauer G A, Malm J R, and Hoffman B F. The P wave and P R interval: Effects of the site of origin of atrial depolarization. *Circulation* 42:653, 1970.
- 11 Waldo A L, Bush H L Jr, Gelband H, Zorn G L Jr, Vitikainen K J, and Hoffman B F. Effects on the canine P wave of discrete lesions in the specialized atrial tracts. *Circ Res* 29:452, 1971.

in New Orleans multiple views of the chest with fluorium in the cephalus revealed marked general sized cardiomegaly (Fig. 1). The electrocardiogram showed low voltage of all complexes and atrial flutter with 2:1 block (Fig. 1). The ventricular rate was 125 beats per minute with an occasional ventricular premature contraction. Digoxin was administered in oral doses to a total of 1.5 mg during the initial 4 days of hospitalization. Quinidine sulphate 200 mg orally every 6 hours was started on the second hospital day. On the fourth hospital day the patient's heart rate was 10 beats per minute. Except for a short period when he had atrial fibrillation which converted to a normal sinus rhythm with a PR interval of 0.23 sec the patient seemed markedly improved. However later the same day as he turned to the left lateral decubitus position for better auscultation of the mitral diastolic rumble he developed ventricular tachycardia and convulsed. He was electrically cardioverted and was transferred to the medical intensive care unit (MICU). Digitalis and quinidine were discontinued. A xylocaine drip of 0.5 mg per minute controlled multifocal ventricular premature contractions. Atrial flutter with 2:1 block recurred.

Chest x-ray the following day showed a large left pleural effusion which yielded 1000 cc of transudative fluid on thoracentesis. A chest x-ray after thoracentesis revealed an infiltrate in the left lower lobe and the patient was started on ampicillin intravenously.

He remained in the MICU with sodium restriction, bed rest, antibiotic and a quiet atmosphere. He did well until the fourteenth hospital day when he developed a temperature of 104°F and rigor. The left pleural effusion had reaccumulated. Kantrex was added to the regimen and 500 cc of transudate were removed by left thoracentesis.

The patient continued to have fever. His urine output diminished, his ascites increased and his neck veins became more engorged. The sputum culture revealed Klebsiella organisms sensitive to Garamycin. Because of a rising blood urea nitrogen and creatinine, Kantrex and Ampicillin were discontinued and he was given a small dose of Garamycin. Intermittent administration of Laix intravenously produced only light diuresis. Digitalis was withheld because of continued ventricular premature contraction.

On the twentieth hospital day he was confused and lethargic. Lumbar puncture was normal except for an opening pressure of 290 mm H₂O due to the high systemic venous pressure. The dosage of Laix was increased to 120 mg intravenously and produced diuresis. Garamycin was administered every other day and the patient became afebrile. A third left thoracentesis removed 1200 cc of transudate.

On the twenty-second day the digitalis medication was raised to 0.5 mg. Second and later transfusions of digitalis. The patient developed short periods of ventricular tachycardia which responded to 100 mg of xylocaine intravenously. On the forty-second hospital day he developed ventricular fibrillation and was defibrillated. The ascites increased and atrial flutter persisted. He finally became unresponsive to the digitalis and various diuretic



Fig. 2 A and B. Immunofluorescent Coxsackie B₁ antibody staining of the aorta of the patient with Coxsackie B₁ viral infection. Panels A and B show stained cytoplasm of cells in the adventitia (Hematoxylin and eosin. Original magnification $\times 125$).

He died on May 17, 1972, 77 days after admission to Charity Hospital in New Orleans.

Methods and materials

Aortic tissue was obtained at autopsy 4 hours after death. The tissue was then stored on ice for 8 hours before fixation in order to assure minimal postmortem changes, although it had previously been shown that the cellular differences between 3 and 24 hours postmortem were not significant in the aorta.⁷ Crossly, the aortic wall had very small scattered whitish spots on the intimal surface; otherwise it was of normal appearance. The tissues were cut to provide longitudinal and cross-sectional orientation and were processed for study by immunofluorescent staining, light microscopic and electron microscopic techniques.

Immunofluorescent staining method. Fresh frozen tissues were cut in sections about 10 μ thick and were fixed in acetone for 10 minutes. After rinsing with phosphate

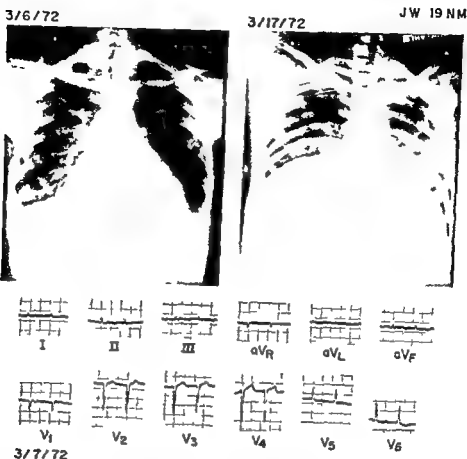
COXSACKIE B₄ VIRAL CARDIOMYOPATHY

Fig. 1 Teleoroentgenograms of the heart and electrocardiogram of the patient with Coxsackie B₄ viral cardiomyopathy at time of admission to Charity Hospital in New Orleans

but no detectable hepatomegaly. The electrocardiogram showed atrial flutter with 2:1 A-V block.

After 10 days of hospitalization and treatment with 20 units of digitalis leaf, there was no improvement. The patient began to have frequent premature ventricular contractions and on March 6, 1972, was transferred to Charity Hospital in New Orleans.

Physical examination on admission to Charity Hospital revealed a very thin and weak Negro man with an arterial blood pressure of 110/90 mm Hg without paradoxical variations. The heart and pulse rates were 136 beats per minute and regular; respiratory rate was 20 per minute and temperature 98.6 F. There was a mild acneiform rash over face and upper back. The neck veins were markedly distended and there were large *a* waves. Few scattered expiratory wheezes and basilar crepitant rales with decreased fremitus, increased dullness, and decreased breath sounds were noted at the bases of both lungs. A right ventricular heave, loud mitral first sound and loud protodiastolic gallop rhythm at the apex were present. A Grade II/VI blowing systolic murmur and a faint diastolic rumble localized to the lower left sternal border in the fourth intercostal space were also present. The murmur did not change with respiration. The abdomen was slightly protuberant and nontender with a fluid wave and shifting dull

ness present. The liver was not enlarged. There was 3+ pitting edema of the feet and legs up to the knees and 3+ post sacral edema. The patient was slightly lethargic with poor attention span.

The laboratory findings on admission were: hemoglobin 13.1 Gm per cent, hematocrit 40.7 per cent, white blood count 8,400 with normal differential, erythrocyte sedimentation rate 4 mm per hour, disulfide preparation negative, lupus erythematosus preparation negative, antistreptolysin O titer 1/50 Todd units, creatine phosphokinase negative, Latex RA negative, fasting blood sugar 78 mg per cent, blood urea nitrogen 14 mg per cent, creatinine 0.7 mg per cent, uric acid 3.6 mg per cent, sodium 131 mEq per liter, potassium 4.8 mEq per liter, chlorine 101 mEq per liter, CO₂ 29 vol per cent, calcium 9.3 mg per cent, phosphorus 4.5 mg per cent, bilirubin 1.6 serum, glutamic oxaloacetic transaminase 130, creatine phosphokinase activity 295, lactic dehydrogenase 560, total serum proteins with A/G ratio 6.7 (3.7/3.0), urinalysis normal with specific gravity of 1.020 and pH of 6.0. Blood was type O and Australian antigen mono screening and febrile agglutinins for Typhoid A, Typhoid O, Paratyphoid B, Proteus OX 19 and Brucella were all negative.

Hospital course. On admission to Charity Hospital



Fig. 4. Photomicrograph of the media of the aorta of the 19 year old patient with Coxsackie B₄ viral infection. Interstitial edema is associated with displacement of the elastic lamellae and smooth muscle cells. (Hematoxylin and eosin. Original magnification $\times 200$.)

hour each. The tissues were embedded in flat molds with appropriate orientation. Polymerization proceeded for 3 days at 60° C. Thick epon sections (0.5 to 10 micron) were cut and stained with toluidine blue in order to ascertain the plane of orientation for selection of the areas to be subsequently sectioned for electron microscopy. Thin sections (500 to 600 Å) from these areas were stained with lead citrate and uranyl acetate and were examined with a Siemens Elmiskop I microscope.

In order to make certain which layer of the aorta was being viewed with the electron microscope when scanning from one layer of the aorta to the next, the tissue blocks were carefully trimmed to specific layers. After thin sections were made from these blocks, an adjacent thick section was prepared to verify that the blocks were limited to the prescribed layers. The thin sections were scanned using the luminal surface as a reference point, thereby estimating the location of a particular cell or cellular layer from that surface. The intimal, medial, and adventitial layers were sectioned in this fashion. In addition, areas were trimmed in which there were no specific pathologic features. The whitish spots observed grossly could not be dis-

Results

Immunofluorescent observations. Positive immunofluorescent staining for Coxsackie B₄ virus antigen was found in the cytoplasm of fibroblasts of the adventitia of the aorta (Fig. 2). This positive immunofluorescent staining was blocked by unlabelled Coxsackie B₄ virus antibody.

Histologic observations. Microscopic observations of the aortic intima, media, and adventitia revealed slight to moderate alterations in the intima and adventitia and slight changes in the media. The endothelial cells were abnormal or were absent in all sections. The intima was fibrotic and contained fat droplets (Fig. 3). In some areas the intima was slightly thickened. Fibroblastic proliferation was present in the subendothelial areas near the lumen of the aorta. The interstices between the proliferating fibroblasts appeared edematous, bumpy, basophilic, and with loosely connected ground substance. In the middle of the thickened intima, fat droplets were accumulated, and the outermost layer of the intima near the media was fibrotic. Cells with small, dense, basophilic nuclei were scattered throughout the intima, indicating a slight degree of inflammatory cell infiltration.

Interstitial edema represented by clear inter spaces between the elastic lamellae and smooth muscle cells associated with

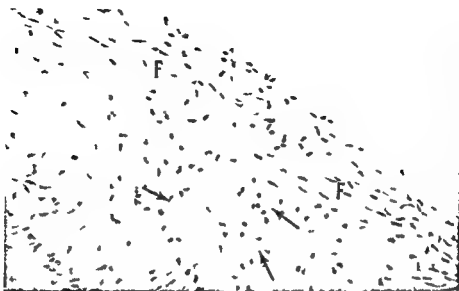


Fig. 7 Photomicrograph of a portion of the thickened intima of the aorta of the 19 year old patient who died of Coxsackie B₄ virus infection. The innermost portion shows fibroblastic proliferation (F) in an edematous background. It is droplets (arras) accumulated within the intima. The outermost portion of the intima is more densely fibrotic than the inner portion. Small dense basophilic nucleated cells, probably inflammatory cells, are scattered throughout the intima (Hematoxylin and eosin. Original magnification $\times 100$).

buffered saline, the sections were stained with fluorescein labelled rabbit antiserum to Coxsackie B₄ virus for 45 minutes in a high humidity chamber. They were then washed again in phosphate buffered saline. The tissues were examined with Cillet Richert UV microscopes.

The antiserum was prepared by infecting rabbits with 0.4 ml Coxsackie virus B₄ antigen intramuscularly and 0.1 ml in the foot pad once a week for 3 weeks. The antigen was obtained by conjugating the purified Coxsackie virus with methylated bovine serum albumin and adding complete Freund's adjuvant.⁸ The rabbits were bled after 3 weeks by cardiac puncture and the serum samples titered. The rabbits were injected once more and exsanguinated one week later. Serum samples were fractionated on a DEAE cellulose ion exchange column and 0.05 molar phosphate buffer elutions (pH 5.0) were collected and labelled with 12.5 micrograms of IITC per milligram of protein in a 10 per cent sodium carbonate bicarbonate buffer (pH 9.5) in a cold environment for 4 hours and then dialyzed against starting buffer with activated charcoal added. Acetone dried tissue powder was used to absorb excess fluorescein and nonspecific antibody prior to staining the tissue sections.

Some sections were tested for immuno-

fluorescent blocking by staining the sections first with unlabelled Coxsackie virus B₄ antibody and then with labelled antibody.

Histologic method Aortic tissue was fixed in 10 per cent neutral formalin, dehydrated and embedded in paraffin. Sections, 7 microns thick, were stained by a routine hematoxylin and eosin method and were examined with a light microscope.

Electron microscopic method Pieces of tissue approximately 1.5 mm by 6 mm long cut through the entire thickness of the aorta were placed in cold 3 per cent glutaraldehyde in phosphate buffer (pH 7.3) for 2 hours. The tissues were subsequently rinsed in 6 changes of fresh phosphate buffer with sucrose (pH 7.3) at 30 minute intervals and were left overnight in cold buffer. Post fixation was with cold 1 per cent osmium tetroxide in phosphate buffer for 15 hours. Dehydration was initiated in a graded series of cold methanol and completed with three changes of pure methanol (20 minutes each) at room temperature. Infiltration was accomplished with pure propylene oxide. The tissues were transferred to fresh vials containing a mixture of $\frac{1}{2}$ propylene oxide and $\frac{1}{2}$ epon and were placed under a lamp overnight. Impregnation with epon was completed with 2 changes of pure epon for 1

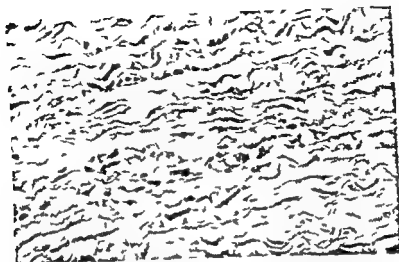


Fig 4 Photomicrograph of the media of the aorta of the 19 year old patient with Coxsackie B₄ viral infection. Interstitial edema is associated with displacement of the elastic lamellae and smooth muscle cell (Hematoxylin and eosin. Original magnification $\times 200$)

hour each. The tissues were embedded in flat molds with appropriate orientation. Polymerization proceeded for 3 days at 60° C. Thick epon sections (0.5 to 1.0 micron) were cut and stained with toluidine blue in order to ascertain the plane of orientation for selection of the areas to be subsequently sectioned for electron microscopy. Thin sections (100 to 600 Å) from these areas were stained with lead citrate and uranyl acetate and were examined with a Siemens Elmiskop I microscope.

In order to make certain which layer of the aorta was being viewed with the electron microscope when scanning from one layer of the aorta to the next the tissue blocks were carefully trimmed to specific layers. After thin sections were made from these blocks an adjacent thick section was prepared to verify that the blocks were limited to the prescribed layers. The thin sections were scanned using the luminal surface as a reference point thereby estimating the location of a particular cell or cellular layer from that surface. The intimal, medial and adventitial layers were sectioned in this fashion. In addition areas were trimmed in which there were no specific pathologic features. The whitish spots observed grossly could not be distinguished in the toluidine blue sec-

Results

Immunofluorescent observations. Positive immunofluorescent staining for Coxsackie B₄ virus antigen was found in the cytoplasm of fibroblasts of the adventitia of the aorta (Fig 2). This positive immunofluorescent staining was blocked by unlabelled Coxsackie B₄ virus antibody.

Histologic observations. Microscopic observations of the aortic intima, media and adventitia revealed slight to moderate alterations in the intima and adventitia and slight changes in the media. The endothelial cells were abnormal or were absent in all sections. The intima was fibrotic and contained fat droplets (Fig 3). In some areas the intima was slightly thickened. Fibroblastic proliferation was present in the subendothelial areas near the lumen of the aorta. The interstices between the proliferating fibroblasts appeared edematous being pale basophilic and with loosely connected ground substance. In the middle of the thickened intima fat droplets were accumulated and the outermost layer of the intima near the media was fibrotic. Cells with small dense basophilic nuclei were scattered throughout the intima indicating a light degree of inflammatory cell infiltration.

Interstitial edema represented by clear interspaces between the elastic lamellae and smooth muscle cells associated with

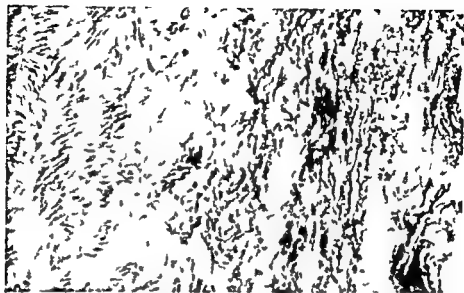


Fig. 5 Photomicrograph showing collagenous tissue proliferation and interstitial hemorrhage in the adventitia of the aorta of the patient (Hematoxylin and eosin. Original magnification $\times 100$)

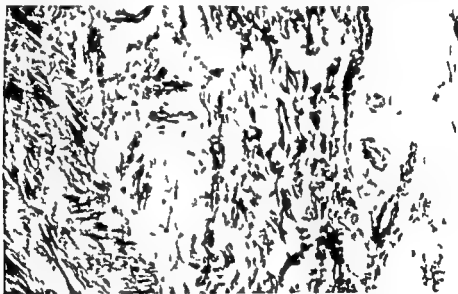


Fig. 6 Photomicrograph of the adventitia of the aorta of the patient. The number of small blood vessels in the adventitia is increased. Collagenous tissue proliferation and interstitial hemorrhage are also present (Hematoxylin and eosin. Original magnification $\times 100$)

disruption of these components was found in some areas of the media (Fig. 4). Proliferation of collagenous tissue was observed in the adventitia (Fig. 5). Interstitial hemorrhage was widespread and varied in severity (Figs. 5 and 6) and the number of small blood vessels in the adventitia was increased (Fig. 6). These changes are indicative of inflammation.

Electron microscopic observations. The endothelial lining of the aorta was absent. This was possibly due to the preparation of the tissue, but the lining may also have been absent prior to the initiation of tissue

preparation in which case it can be speculated that it was pathologically altered and destroyed. In addition there was no organized subendothelial layer. Since no significant alterations were apparent in the media or adventitia by electron microscopy the description of the electron microscopic findings will be limited to the intima of the aorta.

The intima was thickened with the cells scattered and separated by bundles of collagen and widened extracellular spaces. Many of the cells of the intima contained within their cytoplasm varying

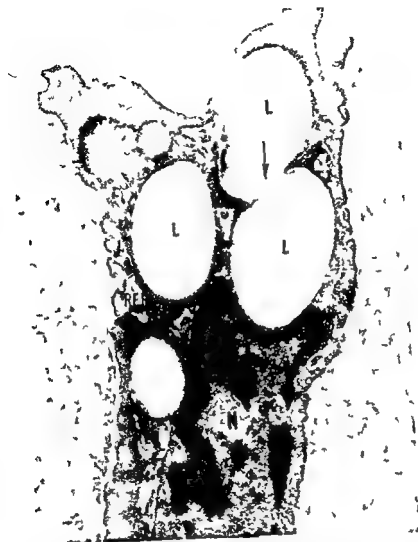


Fig. 7. Electron micrograph of the aortic intima of the patient showing lipid droplets (L) within the cytoplasm of a fibroblast. Two of the droplets have coalesced (arrow). The end of the nucleus (V) is indented by one of the droplets. Clusters of rough endoplasmic reticulum (RER) are apparent. (Original magnification $\times 37,000$)

numbers of lipid droplets which measured 10 to 10 microns in diameter (Fig. 7). In many instances the droplets were coalesced with other droplets often forming chains. Three cell types were observed in which lipid had accumulated. However in some cells the cytoplasm was so full of lipid that the cell type could not be identified with certainty (Fig. 8). The innermost area of the intima closest to the lumen consisted mostly of fibroblast-like cells. However many of the cellular components encountered were so small and fragmentary that their cell type could not be ascertained.

Deeper into the intima approaching the medial layer most of the cells were relatively large smooth muscle cells. Occasionally cells considered to be macrophages were encountered. There was no clearly definable mononuclear infiltration.

Lipid appeared to be deposited mostly in fibroblast-like cells; however the less numerous macrophages contained more lipid per cell. Smooth muscle cells were most frequently devoid of lipid, especially those deeper within the intima. However some smooth muscle cells had 1 or 2 lipid

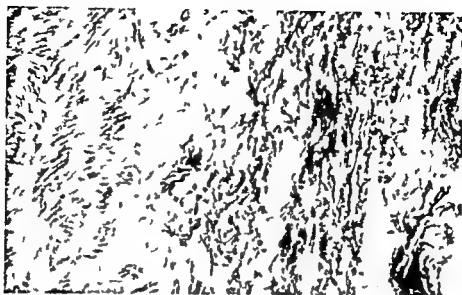


Fig 5 Photomicrograph showing collagenous tissue proliferation and interstitial hemorrhage in the adventitia of the aorta of the patient (Hematoxylin and eosin. Original magnification $\times 100$)



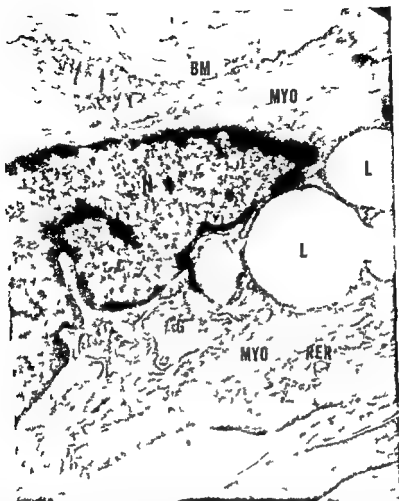
Fig 6 Photomicrograph of the adventitia of the aorta of the patient. The number of small blood vessels in the adventitia is increased. Collagenous tissue proliferation and interstitial hemorrhage are also present (Hematoxylin and eosin. Original magnification $\times 100$)

disruption of these components was found in some areas of the media (Fig 4). Proliferation of collagenous tissue was observed in the adventitia (Fig 5). Interstitial hemorrhage was widespread and varied in severity (Figs 5 and 6) and the number of small blood vessels in the adventitia was increased (Fig 6). These changes are indicative of inflammation.

Electron microscopic observations. The endothelial lining of the aorta was absent. This was possibly due to the preparation of the tissue, but the lining may also have been absent prior to the initiation of tissue

preparation in which case it can be speculated that it was pathologically altered and destroyed. In addition, there was no organized subendothelial layer. Since no significant alterations were apparent in the media or adventitia by electron microscopy, the description of the electron microscopic findings will be limited to the intima of the aorta.

The intima was thickened with the cells scattered and separated by bundles of collagen and widened extracellular spaces. Many of the cells of the intima contained within their cytoplasm varying



Electron micrograph

Fig 9 Electron micrograph of a portion of a smooth muscle cell from the patient's aortic intima. The cell is characterized by the presence of myofilaments (MYO), a basement membrane (BM), and pinocytotic vesicles (arrows). The nucleus (N) with peripheral chromatin is centrally located. Lipid droplets (L) are present in the cytoplasm. A Golgi apparatus of several laminate cisternae (G) can be seen close to the nucleus. Elements of the rough endoplasmic reticulum (RER) are also present. (Original magnification $\times 27,000$)

filaments within the cytoplasm, presence of pinocytotic vesicles, and presence of a surrounding basement membrane. The smooth muscle cells had a well-developed Golgi apparatus. Rough endoplasmic reticulum was common, though not abundant. Most of the smooth muscle cells had only one or two small lipid droplets or were without lipid. Occasionally a cell was encountered in which several lipid droplets were apparent (Fig. 9).

The third type of cell in which lipid was found was considered to be a macrophage (Fig. 8). This cell may also represent the foam cell previously described in aortic

fatty streaks.¹² In the cell illustrated in Fig. 8 the nucleus was displaced to one side by the large amount of accumulated lipid. The cytoplasm was condensed in areas as a result of crowding by the lipid. Coalescence of lipid droplets was a common observation. Mitochondria were scarce and rough endoplasmic reticulum was sparse. Occasional dense bodies, which are possibly cytolysomes, were seen. Vesicles smaller than the lipid droplets and with electron dense contents were also seen. Small thin projections of the cytoplasm, which protruded into the extracellular space, were frequently observed in these

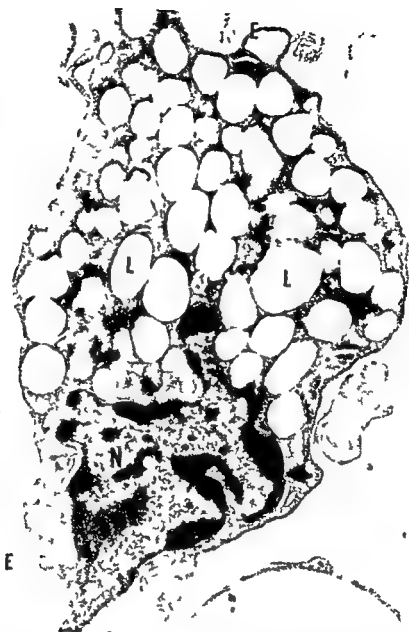


Fig. 8 Electron micrograph of a macrophage from the intima of the patient's aorta. The cytoplasm contains an abundance of lipid droplets of various sizes (L). The nucleus (N) is displaced to one side by crowding of lipid. The cytoplasm is dense due to compression. Small thin protoplasmic extensions (E) protrude from the cell periphery. (Original magnification $\times 21,600$.)

droplets, and a few had more than 2 droplets (Fig. 9).

The fibroblast-like cells (Fig. 7) were identified by their elongated profile and irregular projections into the extracellular space, absence of myofibrils and absence of a surrounding basement membrane. The cell demonstrated in Fig. 7 represents a protoplasmic extension of a fibroblast containing several lipid droplets. The cytoplasm is homogeneously dense. The nucleus is elongated, with peripheral chromatin. Short profiles of rough endo-

plasmic reticulum can be seen. Pinocytotic vesicles were not apparent. Thin fibrils arranged loosely into small bundles were found within the extracellular spaces. Extracellular fibrous material was observed in close apposition to the plasma membrane.

The smooth muscle cells were generally larger than the fibroblast-like cells and were disposed more frequently near the media. The smooth muscle cells (Fig. 9) were distinguishable from the fibroblast-like cells (Fig. 7) by the presence of myo-

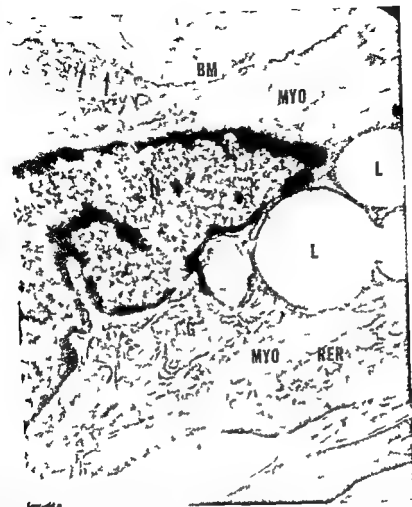


Fig. 9. Electron micrograph of a portion of a smooth muscle cell from the patient's aortic intima. The cell is characterized by the presence of myofilaments (MYO), a basement membrane (BM), and pinocytotic vesicles (arrows). The nucleus (N) with peripheral chromatin is centrally located. Lipid droplets (L) are present in the cytoplasm. A Golgi apparatus of several laminate cisternae (G) can be seen close to the nucleus. Elements of the rough endoplasmic reticulum (RER) are also present. (Original magnification $\times 27,000$.)

filaments within the cytoplasm, presence of pinocytotic vesicles, and presence of a surrounding basement membrane. The smooth muscle cells had a well developed Golgi apparatus. Rough endoplasmic reticulum was common though not abundant. Most of the smooth muscle cells had only one or two small lipid droplets or were without lipid. Occasionally a cell was encountered in which several lipid droplets were apparent (Fig. 9).

The third type of cell in which lipid was found was considered to be a macrophage (Fig. 8). This cell may also represent the foam cell previously described in aortic

fatty streaks.¹⁷ In the cell illustrated in Fig. 8 the nucleus was displaced to one side by the large amount of accumulated lipid. The cytoplasm was condensed in areas as a result of crowding by the lipid. Coalescence of lipid droplets was a common observation. Mitochondria were scarce and rough endoplasmic reticulum was sparse. Occasional dense bodies which are possibly cytolysosomes were seen. Vesicles smaller than the lipid droplets and with electron dense contents were also seen. Small thin projections of the cytoplasm which protruded into the extracellular space were frequently observed in these

cells. The macrophage clearly lacked a basement membrane.

Discussion

The alterations observed in this patient are indicative of inflammation of the aorta. These include thickening, fibroblastic proliferation, fibrosis and basophilic infiltration of the intima, edema of the media, accumulation of lipid, proliferation of collagenous tissue, hemorrhage and increased vascularization in the adventitia. The important finding of fairly extensive fat deposition within the intima is suggestive of atherosclerotic changes in this young man who died of a viral infection. Furthermore, immunofluorescent antibody staining was positive for Coxsackie B₄ virus in cells of the aorta. Because of the young age of this patient (19 years) the possibility that the lipid deposition resulted only from a progressive aging phenomenon is unlikely. Also from the clinical data it is unlikely that the lipid deposits were due to either hypertension or to a high cholesterol diet in this patient.

Aortitis indicated by inflammatory changes in the aortic wall has been observed in syphilis,⁹ rheumatic fever,¹⁰ rheumatoid arthritis¹¹ and pulseless disease¹² or aortitis syndrome.¹³ Fibrous thickening of the intima and the adventitia and disruption and loss of elastic and muscular fibers in the media are the common morphologic alterations of the aortic wall in these diseases.

Lipid deposition on the other hand which has been considered an early manifestation of atherosclerotic lesions⁷ and which is usually not observed in vasculitis was observed in this patient. It is evident, therefore, that there were changes in the aorta associated with early atherosclerotic changes. It is not possible to decide from our present data which of these changes occurred first. But, when it is realized that this patient was young and that immunofluorescent antibody staining for Coxsackie B₄ virus antigen was positive in the aorta it should be considered that a viral infection of the aorta at least might have predisposed to the lipid deposition. Deposition of lipid in injured tissue can occur extremely rapidly in a few hours to a few days. The fact that positive immunofluorescent anti-

body staining for antigen of Coxsackie virus B₄ was observed in the cells of the adventitia of the aorta of this patient suggests the possibility that the inflammatory changes in the aortic wall were caused by viral infection. Upon examination with the electron microscope, it was found that most of the intimal cells contained lipid in varying quantities. However, no extracellular lipid which has been considered an early manifestation of the formation of atherosclerotic lesions,⁷ was found.

Most of the lipid containing cells could be recognized as fibroblast like cells. They were usually closest to the luminal surface of the aorta. Lipid containing smooth muscle cells were more frequently encountered and lipid containing fibroblasts were less frequently encountered in areas progressively more distal to the luminal surface. The smooth muscle cells became increasingly surrounded by elastin fibers in the more distal regions. Many of the deeper smooth muscle cells were without lipid, although, as indicated above, some contained 1 or 2 droplets and a few contained numerous droplets of lipid. These observations are contrary to earlier reports^{4,7} of atherosclerotic lesions, both proliferative lesions (non necrotic) and fatty streaks. In those reports the predominant cells were mature smooth muscle cells and the lipid laden cells were either monocytes,⁴ smooth muscle cells,⁶ macrophages⁴ or foam cells.^{5,7} Only a few cells which could be considered as either macrophages or foam cells were encountered in the aorta of the patient described here. No cells could be recognized as monocytes.

In man fat droplets appear in smooth muscle cells at an early stage in the formation of atherosclerotic lesions.¹⁴ As the accumulation of fat continues the smooth muscle cells develop into myogenic foam cells.¹⁴ The foam cells subsequently degenerate and empty their contents into the extracellular space. This fat in turn, is phagocytosed by macrophages which in time may assume the features of foam cells.¹⁴ In a recent report it was indicated that some foam cells may be derived from smooth muscle cells whereas others, which resemble macrophages appear to be derived from monocytes previously circulating in the blood stream.⁶ It has not been

1 m 86
mb 4

established whether the macrophages are altered primitive cells which migrate from the aortic media or whether they are derived from the circulating blood and gain entrance into the intima after migrating through the endothelial lining of the aorta.

It should be pointed out that lesions compatible with early atherosclerotic changes have been described in our patient. It is proposed that the changes reported in this patient were a result directly or indirectly of the accompanying Coxsackie B₁ viral infection. It seems possible that injury to the arterial vessels by a viral agent could be one of the factors which initiate a site of injury which through time arterial blood pressure and lipoproteins and cholesterol circulating in the blood is repaired resulting in the formation of scars and other atherosclerotic changes observed in all people especially later in life. Arteriosclerosis has a beginning; it starts in young people and accumulates other injuries with time to become more extensive with age. Viremia with viral injury of the arterial system must be seriously considered as an initiating factor of atherosclerosis. Viral infections begin with life and continue throughout life. We have shown in previous studies in experimental animals that viruses can injure extensively the aorta, coronary arteries and other arteries of mice.¹⁴ It is our contention that atherosclerosis and arteriosclerosis with calcification and ulceration are late scars and reparative phenomena of earlier and more fundamental lesions of the arterial vessels.¹⁷ This concept would also explain the reasons for the particular sites of the atherosclerotic and arteriosclerotic lesions noted by pathologists and clinicians.

Summary

A 19 year old young man who died of viral cardiomyopathy had positive staining of the myocardium, kidney, pancreas and aorta with immunofluorescent antibodies for Coxsackie B₁ virus. The aorta was found to have an accumulation of lipid material in cells of the intima. The lipid deposits were located electron microscopically in fibroblasts, smooth muscle cells and macrophages. It is suggested that viral infections so common in man may produce local sites

scars and lipid, calcium and other deposits which represent the well known lesions of atherosclerosis and arteriosclerosis. These scars and late lesions must have a beginning and from the manifestations of immunofluorescent antibodies, histopathologic and electron microscopic findings in the aorta of the patient described here as well as from the clinical data, it is suggested that Coxsackie B₁ virus may initiate atherosclerosis and arteriosclerosis in man.

REFERENCES

1. McGill H C, Frink M H and Geer J C. Aortic lesions in hypertensive monkeys. *Arch Pathol* 71:96 1961.
2. Armstrong M L, Warner E D and Conner W E. Regression of coronary atheromatosis in the rhesus monkeys. *Circ Res* 27:49 1970.
3. Bortz W M. Reversibility of atherosclerosis in cholesterol fed rabbits. *Circ Res* 22:135 1968.
4. Tucker C F, Catsulis C, Strong J P and Eggen D A. Regression of early cholesterol induced aortic lesions in rhesus monkeys. *Am J Pathol* 65:493 1971.
5. Peterson M, Day A J, Tume R H and Eisenberg E. Ultrastructure, fatty acid content and metabolic activity of foam cells and other fractions separated from rabbit atherosclerotic lesions. *Exp Molec Pathol* 15:157 1971.
6. Scott R F, Jones R, Daoud A S, Zumbo O, Coulton F and Thomas W A. Experimental atherosclerosis in rhesus monkeys. II. Cellular elements of proliferative lesions and possible role of cytoplasmic degeneration in pathogenesis as studied by electron microscopy. *Exp Molec Pathol* 7:44 1967.
7. Geer J C. Fine structure of human aortic intimal thickening and fatty streaks. *Lab Invest* 14:164 1965.
8. Williams C A and Chase M W. Methods in immunology and immunochemistry, vol 1. New York: 196. Academic Press, pp 204-205.
9. Heggqvist H A. Syphilitic aortitis: a clinicopathologic autopsy study of 100 cases 1950 to 1960. *Circulation* 29:346 1964.
10. Fappenheimer A M and von Glahn W C. Lesions of the aorta associated with acute rheumatic fever and with chronic cardiac disease of rheumatic origin. *J Med Res* 44:489 1924.
11. Clark W S, Kulka J J and Bauer W. Rheumatoid aortitis with aortic regurgitation: an unusual manifestation of rheumatoid arthritis (including polydactylitis). *Am J Med* 22:580 1957.
12. Niu T. Pathology of pulseless disease: a systematic study and critical review of twenty one autopsy cases. *Angiology* 14:225 1961.
13. Ueda H et al. Clinical and pathological studies of aortitis syndrome: committee report. *Jap Heart J* 9:16 1968.

- 14 Haust M D More R H Bencoame S A and Balis J U Electron microscopic studies in human atherosclerosis Extracellular elements in aortic dots and streaks *Exp Mol Pathol* 6:300 1967
- 15 Sohal R S Burch G E Chu K C Leiderman E and Colcolough H L Ultrastructural changes in cardiac capillaries of coxsackie virus B₁ infected mice *Lab Invest* 19:399 1968
- 16 Burch G E Tsui C Y and Harb J M Pathologic changes of aorta and coronary arteries of mice infected with coxsackie B₁ virus *Proc Soc Exp Biol Med* 137:657 1971
- 17 Burch G E The etiology of arteriosclerosis-A thought *AM HEART J* 83:434 1972

Charcot-Marie-Tooth disease, Wolff-Parkinson-White syndrome, and abnormal intracardiac conduction

Dorrance Bowers M D
Kelowna B C Canada

This report describes the occurrence of the Wolff Parkinson White syndrome (anomalous atrioventricular excitation and recurrent paroxysmal tachycardia) in a young man with Charcot Marie Tooth disease (peroneal muscular atrophy). In addition to the fascinating compounding of eponyms this patient is of interest because his electrocardiograms on occasion showed delayed conduction from the sino auricular node to the ventricles both via the normal path and via the accessory path and also left anterior fascicular block.

Case report

This Canadian born white male had been known to have paroxysmal supraventricular tachycardia (Fig 1) since early life with intermittent normal ventricular excitation (Fig 2) and intermittent anomalous ventricular excitation (Fig 3).

At the age of 22 years he was noted to display the features of peroneal muscular atrophy namely wasting of the distal muscles of the extremities, high arched feet and curled toes. The conduction velocity along his left median nerve was measured at 49 meter per second and along his left ulnar nerve at 33 meters per second these values are at the lower limit of normal and are quite unusual for a healthy young man.

This patient's electrocardiograms intermittently showed delayed atrioventricular conduction both by the accessory path (Figs 3 and 4) and by the normal path (Fig 4). This atrioventricular conduction delay occurred in the absence of digitalis administration and was not modified by the resumption of

the upright posture or by the intravenous administration of atropine 1.2 mg. The patient's electrocardiograms also showed left anterior fascicular block.

There was no knowledge of the occurrence of the Wolff Parkinson White syndrome or of Charcot Marie Tooth disease in other members of his family. His parents, his three younger brothers and his only sister were examined for clinical evidence of peroneal muscular atrophy. No evidence of this disorder was found. Because his siblings ranged in age from 10 to 21 years when examined it remains possible that they may manifest the stigmata of peroneal muscular atrophy in the future. Electrocardiograms from his parents and four siblings showed no conduction abnormalities or tachyarrhythmias. Because the Wolff Parkinson White phenomenon is commonly intermittent these normal electrocardiograms do not exclude the possibility of intermittent anomalous atrioventricular excitation in these other members of his family.

Discussion

This patient displayed the classical clinical features of peroneal muscular atrophy; the diagnosis was supported by the slow rate of conduction along his peripheral nerves. He displayed also the classical clinical features of the Wolff Parkinson White syndrome namely anomalous atrioventricular excitation and recurrent paroxysmal supraventricular tachycardia. His electrocardiograms were unusual in that impulses conducted to the ventricles both by the normal path and by an accessory path at times showed prolonged atrioven-

From the Department of Medicine, Kelowna General Hospital, Kelowna, B. C. Canada.
Received for publication April 10, 1973.

Reprint requests: Dr. Dorrance Bowers, M.D., 1737 Broadway St., Kelowna, B. C. Canada.



Fig 1 Regular supraventricular tachycardia with ventricular rate approximately 187 per minute

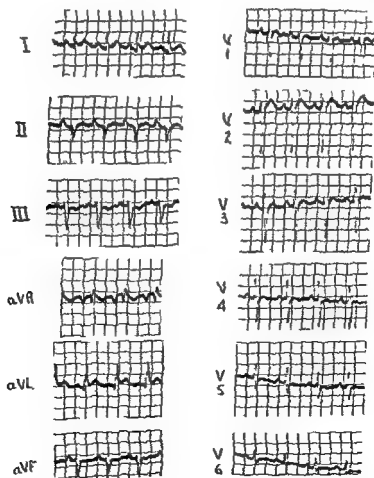


Fig 2 Sinus rhythm with atrioventricular conduction via the normal path (PR interval \approx 0.20 sec). The QRS complexes are similar to those in Fig 1 indicating that during that paroxysm of supraventricular tachycardia ventricular excitation occurred via the normal path. Note also the left anterior fascicular block.

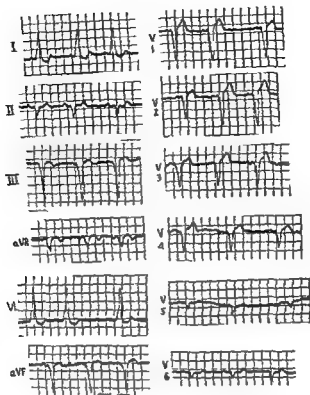


Fig. 3 Sinus arrhythmia with anomalous atrioventricular excitation. The P delta intervals vary from 0.15 to 0.21 second in duration. According to the criteria of Rosenbaum and associates² this electrocardiogram is an example of the Wolff Parkinson White syndrome (Type B). Bruyneel⁶ has proposed that the initial delta vector be the criterion for the electrocardiographic subdivision of the WPW syndrome regardless of the duration and sense of the main QRS vector—a positive delta wave in Lead V_1 indicating Type A and a negative delta wave in Lead V_1 indicating Type B. If this criterion is employed this electrocardiogram must be classed as an example of the WPW syndrome (Type A).

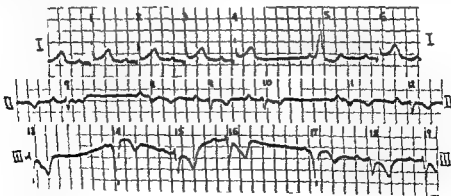


Fig. 4 Intermittent anomalous atrioventricular excitation with delayed AV conduction both by the normal path ($PQ = 0.24$ second) and via the accessory route ($P\delta = 0.17$ second). In complexes 7, 10, 13, and 16 the e wave thought to be retrograde premature P waves caused by reentry of impulses into the atrium retrograde direction via the accessory atrioventricular path. Because the P waves are negative in Leads II and III Wolff Parkinson White syndrome Type A (resulting from a bypass connecting the left atrium and left ventricle) is suggested. Note that the intermittent single anomalous atrioventricular excitation complexes (5, 8, 11, 14, and 17) occur after long P-R intervals presumably due to resetting of the SA node by the retrograde P waves. The absence of delta waves after shorter P-R intervals suggests that the refractory period of the accessory path is longer than that of the normal path.

Table 1 Reported patients with peroneal muscular atrophy and abnormal intracardiac conduction

| Author | Patient | Peroneal muscular atrophy | AV conduction abnormality | Intraventricular conduction abnormality | Arrhythmia |
|----------------------------|---|---------------------------|---|---|---|
| Legrand et al ¹ | Male aged 44 | Variant | IR = 0.30 sec | Left bundle branch block | None |
| Leak ² | Male aged 39 | Typical | Varying AV block in association with atrial flutter | None | Atrial flutter |
| Littler ³ | II 1 male aged 41 | Typical | Complete AV block with Stokes-Adams attacks | Previous ECG showed right bundle branch block and left axis deviation | None |
| | I 2 male aged 74 | Bilateral pes cavus | None | Right bundle branch block and left axis deviation | None |
| | II 2 female aged 40 | Bilateral pes cavus | None | Incomplete right bundle branch block | None |
| Hay et al ⁴ | Same patient as Littler ³ II 1 | | | | |
| Present report | Male aged 33 | Typical | Intermittent first degree AV block | Left anterior fascicular block | W P W syndrome (type 1) paroxysmal supraventricular tachycardia |

tricular conduction and also left anterior fascicular block.

It is tempting to speculate regarding the significance of intracardiac conduction disturbances in patients with peroneal muscular atrophy. The possibility that this association is merely coincidental seems unlikely in view of the increasing number of patients in whom this association has been recognized (Table 1). Littler³ has reviewed in detail possible genetic explanations which might account for the association of cardiac conduction defects and peroneal muscular atrophy in the family he reported. Surely the possibility must also be considered that the prolonged peripheral nerve conduction and prolonged intracardiac conduction in these patients have a common etiology, although peripheral nerve conduction is based on nerve cells and cardiac conduction on specialized myocardial cells; this anatomic difference need not imply a physiologic difference in the electrochemical transmission of impulses via specialized myocardial cells as opposed to nerve cells.

Summary

A patient with Charcot Marie Tooth disease, Wolff Parkinson White syndrome

and abnormal intracardiac conduction has been reported.

The peripheral nerve conduction velocity studies were carried out by Dr W St J Buckler Vancouver B C Canada. The author is grateful to Dr F C MacInnes Summerland B C Canada for referring this patient and to Dr Alfred Pick of Michael Reese Hospital Chicago Illinois for his comments on the patient's electrocardiograms.

REFERENCES

- 1 Legrand R, De ruelles J and Minouvier A. Troubles cardiaques graves dans un cas d'amyotrophie de Charcot Marie avec ataxie. Arch Mal Coeur 17:154 1950.
- 2 Leak D. Paroxysmal atrial flutter in peroneal muscular atrophy. Br Heart J 23:376 1961.
- 3 Littler W A. Heart block and peroneal muscular atrophy. A family study. Q J Med 39:431 1970.
- 4 Kay J M, Littler W A and Meade J B. Ultrastructure of myocardium in familial heart block and peroneal muscular atrophy. Br Heart J 34:1081 1972.
- 5 Rosenbaum I F, Hecht H H, Wilson F N and Johnston I D. The potential variation of the thorax and esophagus in anomalous atrioventricular excitation (Wolff Parkinson White syndrome). Am Heart J 29:781 1945.
- 6 Bruynel K J J. Wolff Parkinson White syndrome (Correspondence). Circulation 47:433 1973.
- 7 Schamroth I. How to approach an arrhythmia. Circulation 47:470 1973.

Table 1 Reported patients with peroneal muscular atrophy and abnormal intracardiac conduction

| Author | Patient | Peroneal muscular atrophy | IV conduction abnormality | Intracardiac conduction abnormality | Arrhythmia |
|----------------------------|------------------------------|---------------------------|---|---|--|
| Légrand et al ¹ | Male aged 41 | Variant | PR = 0.30 sec | Left bundle branch block | None |
| Leak ² | Male aged 39 | Typical | Varying AV block in association with atrial flutter | None | Atrial flutter |
| Littler ³ | II 1 male aged 41 | Typical | Complete AV block with Stokes-Adams attacks | Previous ECG showed right bundle branch block and left axis deviation | None |
| | II 2 male aged 44 | Bilateral pes cavus | None | Right bundle branch block and left axis deviation | None |
| | II 2 female aged 10 | Bilateral pes cavus | None | Incomplete right bundle branch block | None |
| Kay et al ⁴ | Same patient as Littler II 1 | | | | |
| Present report | Male aged 12 ⁵ | Typical | Intermittent first degree AV block | Left anterior fascicular block | W P W syndrome (type A) paroxysmal supra ventricular tachycardia |

tricular conduction and also left anterior fascicular block.

It is tempting to speculate regarding the significance of intracardiac conduction disturbances in patients with peroneal muscular atrophy. The possibility that this association is merely coincidental seems unlikely in view of the increasing number of patients in whom this association has been recognized (Table 1). Littler³ has reviewed in detail possible genetic explanations which might account for the association of cardiac conduction defects and peroneal muscular atrophy in the family he reported. Surely the possibility must also be considered that the prolonged peripheral nerve conduction and prolonged intracardiac conduction in these patients have a common etiology, although peripheral nerve conduction is based on nerve cells and cardiac conduction on specialized myocardial cells. This anatomic difference does not imply a physiologic difference in the electrochemical transmission of impulses via specialized myocardial cells as opposed to nerve cells.

Summary

A patient with Charcot Marie Tooth disease, Wolff Parkinson White syndrome

and abnormal intracardiac conduction has been reported.

The peripheral nerve conduction velocity studies were carried out by Dr W. St. J. Buckler, Vancouver, B. C., Canada. The author is grateful to Dr F. C. MacInnes, Summerland, B. C., Canada, for referring this patient and to Dr Alfred Pick of Michael Reese Hospital, Chicago, Illinois, for his comments on the patient's electrocardiogram.

REFERENCES

1. Légrand R, De ruelles J, and Minouvier A. Troubles cardiaques graves dans un cas d'amyotrophie de Charcot-Marie avec ataxie. Arch Mal Coeur 13:154, 1950.
2. Leak D. Paroxysmal atrial flutter in peroneal muscular atrophy. Br Heart J 23:326, 1961.
3. Littler W. A. Heart block and peroneal muscular atrophy. A family study. Q J Med 19:431, 1970.
4. Kay J, M. Littler W. A. and Meade J. B. Ultrastructure of myocardium in familial heart block and peroneal muscular atrophy. Br Heart J 34:1081, 1972.
5. Roenigum J. I., Hecht H. H., Wilson J. N., and Johnston I. D. The potential variation of the thorax and esophagus in idiopathic intracardiac excitation (Wolff Parkinson White syndrome). Am Heart J 29:281, 1945.
6. Bruyneel K. J. J. Wolff Parkinson White syndrome (Correspondence). Circulation 47:433, 1973.
7. Schramm I. How to approach an arrhythmia. Circulation 47:420, 1973.

though serial ECG's and enzymes were not diagnostic. Five days later the patient was discharged from the coronary surveillance unit. On the seventh hospital day seven days after the presumed infarct a pericardial friction rub was heard with a low grade fever which then disappeared spontaneously. On the twenty-fifth hospital day a holosystolic murmur was first heard in the apex with radiation to the axilla. The heart tones were noted to be muffled and distant. The patient however was improving clinically. There was no evidence of congestive heart failure and chest pains were less frequent although the ECG's continued to show a possible active myocardial or pericardial process. On the tenth and twenty-first hospital days LE preparations were positive and on the thirtieth hospital day the B C was 72 mg per 100 ml. On the twenty-fifth hospital day azathioprine was started because of the possible activity of the disease and in an effort to be able to reduce the steroid dose of 30 mg prednisone daily. By the thirtieth hospital day the patient was completely asymptomatic. The renal status was stable and he was gradually ambulating without any problems. On the thirty-third hospital day the patient was discharged.

Two days after discharge from the hospital the patient died.

Discussion

DR IZZY SOMMERS (MODERATOR) The case for discussion today is that of a 25 year old man who presented with systemic lupus erythematosus (SLE) and the nephrotic syndrome. We are privileged to have with us to solve the problems of this case Dr Norman M Simon Associate Professor of Medicine at Northwestern University Medical School.

DR N M SIMON The case presents an unusual challenge in that the diagnosis of SLE appears to be well established. The course is an extended one and my task is to sort out the nature and extent of the systems involved by the SLE process and to determine whether other significant diseases also were present.

The patient was a 25 year-old Japanese man. In our experience at Northwestern University Medical Center we have noted an increased prevalence of people of the Oriental race in patients with SLE.¹ He was found to have SLE at the age of 4 years and was started on corticosteroid therapy. Two years later he first showed renal involvement manifested by the nephrotic syndrome. Subsequently he developed a number of serious complications related to steroids which were discontinued after a treatment period of eight years. The first renal biopsy was performed nine years

after the onset of renal disease and the tissue was reported to show diffuse membranous glomerulonephritis. This is one type of renal lesion found in SLE. Involvement of the kidneys is common in lupus occurring in 50 to 80 per cent of patients generally early in the clinical course. The renal lesions of SLE have been well defined by Pollak and Pirani² and Baldwin and co-workers.³ In addition to membranous glomerulonephritis these include a mild alteration labeled as minimal glomerular involvement or focal glomerulonephritis and a severe abnormality described as active or diffuse glomerulonephritis. Each lesion appears to have its own natural history and there is little tendency for one to progress into another.

The patient was then started on treatment with azathioprine which was continued for several years until the disease was inactive. He returned a year or two later with signs of acute occlusion of the major veins of the left leg extending into the inferior vena cava. The occurrence of right flank pain suggests involvement of the renal veins as well which might account for the observed increase in proteinuria. Indeed it is possible that renal vein thrombosis may have been the underlying disorder from the very beginning. The renal biopsy finding of membranous glomerular changes is compatible with this speculation.

However there are alternative explanations to account for thrombotic venous disease in this patient. First SLE is a vasculitis which may affect veins as well as arteries. Second significant abnormalities in blood coagulation are found in patients with the nephrotic syndrome regardless of etiology.⁴ These include increased levels of plasma fibrinogen factors V combined VII and VIII and platelets and accelerated thromboplastin generation. The coagulation abnormalities disappear as the nephrotic syndrome goes into remission. In section formerly was a major threat to nephrotic patients before the introduction of antibiotics thromboembolism now stands as a serious complication of their disease.

The patient was started on treatment with oral anticoagulants but required readmission to the hospital a short time later with evidence of active disease. A large

was reinstituted and prednisone was continued at 30 mg daily. Coumadin, prednisone and quinidine were continued after discharge.

In July 1971 he was admitted for the fifth and last time with a two-day history of progressive shortness of breath, increasing leg swelling, fever, tachycardia and coughing. He denied any chest pain or hemoptysis and there was no abnormal bleeding. His cough was productive of whitish sputum. He also complained of pain in his testicles for about three days.

Physical examination on admission. The patient appeared Cushingoid and was in moderate respiratory distress. Vital signs included blood pressure of 116/80 mm Hg, pulse 110 per minute with occasional premature beats, respirations 40 per minute and labored and temperature was 102° F. The skin was warm with good turgor. There was a dry, scaly, slightly red maculopapular rash over the face with butterfly distribution and well delineated borders. Similar lesions appeared on the dorsum of both hands and in the groin areas with strikingly well delineated borders. Acneiform eruptions were present in the neck, chest and back. The head was normocephalic. The eyes showed bilateral iridectomy scars and absent left lens. The fundi showed flat discs with arteriolar narrowing and A-V nicking, no hemorrhages, exudates or cytoid bodies were seen. The neck was supple with moderate jugular venous distention at 45°. The hepatjugular reflux was positive. The chest expanded symmetrically without splinting; there was diffuse wheezing with bilateral basilar rales. There was no evidence of consolidation or effusion. The point of maximum cardiac impulse was not palpable or visible. The heart tones were of good quality. S₁ and S₂ were normal. S₂ was intermittently heard at the apex and S₄ was present. No murmurs or rubs were heard. The abdomen was distended; no fluid waves could be elicited. There were numerous venous collaterals seen on the abdominal wall. The liver span was 12 cm with a smooth, nontender edge. The spleen was not felt. There was bilateral pedal edema. There was no evidence of thrombophlebitis. Homan's sign was negative. There was no clubbing. The nails were pink, white and scaly. The right epididymis was markedly swollen and inflamed and extremely tender to palpation. The neurological examination revealed no abnormalities.

Laboratory data. The hemoglobin was 14.5 Gm per cent, hematocrit 44.7 per cent and RBC count 4.5 million per cubic millimeter. WBC count was 8,600 per cubic millimeter with a differential of 78 neutrophils, 1 band, 15 lymphocytes, 0 monocytes and numerous platelets. Urinalysis showed a pH of 6, specific gravity 1.005, a large amount of protein and negative for sugar and acetone. There were seven to fifteen RBCs and two to five WBCs per high power field. Also the urinalysis showed a few bacteria and was positive for doubly refractile fat bodies. The chest x-ray showed cardiomegaly with pulmonary vascular congestion. An ECG (electrocardiogram) showed old posteroinferior myocardial infarction with frequent PVCs (premature ventricular contractions). Sodium was 148, potassium 4.3, chloride 95 and CO₂ 23 mEq per liter. BUN was 42, creatinine 1.7 mg per cent. The arte-

rial blood gases showed a pH of 7.48, a Pco₂ of 76 and a Po₂ of 139 mm Hg while the patient was receiving oxygen by nasal cannula. The SGOT (serum glutamic oxalacetic transaminase) was 35, SGPT (serum glutamic pyruvic transaminase) 42, LDH (lactic dehydrogenase) 357, CPK (creatinine phosphokinase) 7 units. Serial examinations of these enzymes did not show any significant increase. The alkaline phosphatase was 2.5 Bodansky units. The total bilirubin was 0.3, direct 0.2 mg per 100 ml. Total protein was 5.3 Gm per cent with albumin of 2.4 Gm. The serum cholesterol was 220 and triglycerides 89 mg per 100 ml. Serum complement done on July 23, 1971 was 80 mg per 100 ml on July 29, 1971 120 mg per 100 ml. The IgG was 1,000, IgA 210, IgM 350 mg per 100 ml. Cryoglobulin was present. A 24-hour urine protein showed 2.23 Gm per 24 hours. The RA factor was negative. The first two IE preparations were negative but later these became positive. The prothrombin time was 22.8 seconds, PTT was 46.6 seconds with a control of 39.0. Blood and sputum cultures for acid fast bacilli were negative. A urine culture was also negative as well as stool cultures. Skin lesions on the face, hands and groin showed mycelium and pores on the potassium hydroxide smears.

Hospital course. The patient was extremely anxious and acutely short of breath. He was thought to be in congestive heart failure and treatment was begun with diuretics, digitalis and sedation with good response. The possibility of pulmonary embolization was strongly suspected; heparin therapy was started. An emergency lung scan showed evidence only of pulmonary congestion and other nonspecific findings. Cardiology and cardiovascular consultation suggested that conservative management with out further diagnostic studies was indicated. By evening the patient had markedly improved and complained only of pains in his right testicle which subsequently subsided with analgesics, tetracycline and local heat. The temperature was elevated for two days but came down with improvement of the epididymitis. On the second hospital day while being weighed he developed sudden severe substernal chest pain with shortness of breath, diaphoresis and transient hypotension. Physical examination was unrevealing and the ECGs remained unchanged. He responded readily to oxygen and morphine. He was then kept at complete bed rest and heparin, prednisone and quinidine were continued. He then stabilized. Dermatology consultants diagnosed fungal infections of the skin and nails by potassium hydroxide smears. Therapy with Tinctin was followed by marked improvement. Meanwhile renal function was slightly deteriorating with creatinine of 3 and BUN of 80 mg per 100 ml. The proteinuria, however, at most was 2.5 mg per 24 hours.

On the seventh hospital day episodes of minor chest pain again occurred with shortness of breath. Serial ECGs showed persistent elevation of ST segments suggestive of an active myocardial and/or pericardial process. On the tenth hospital day more severe chest pain occurred with shortness of breath, sweating, bradycardia and hypotension. The patient was transferred to the coronary surveillance unit with a diagnosis of acute myocardial infarction. Al-

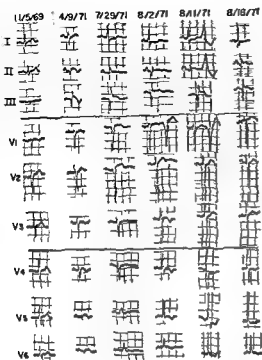


Fig 1 ECGs taken on Nov 5 1969 April 9 July 29 Aug 11 and Aug 18 1971 (see text)

The chest film at the time of the first admission showed the heart size to be normal. The first phlebogram on Aug 18 1969 (Fig 2) showed almost complete occlusion by a large thrombus in the left femoral vein and this is seen to extend upward into the vena cava with occlusion of the vena cava at the level of the fourth lumbar vertebra. There is extensive collateralization of the veins in the perivertebral plexus. There are small clots in the right femoral vein. The examination was repeated about three months later on Nov 5 1969 (Fig 3). It is evident that the clots in the right femoral vein have disappeared but there is essentially no difference in the left femoral vein thrombus. There continue to be extensive venous collaterals. Chest films showed that the patient went on to develop cardiac enlargement and intrapulmonary effusion both of which improved in response to treatment prior to the patient's death.

DR SIMON: Were there signs of occlusion of the renal veins?

DR LEVIN: I cannot tell whether or not



Fig 2 Phlebogram of Aug 18 1969 showing occlusion of left femoral vein and of inferior vena cava at the level of the fourth lumbar vertebra. There are small thrombi in the right femoral vein and extensive collateral veins.



Fig 3 Phlebogram of Nov 5 1969. The thrombi in the right femoral vein are no longer present. The status of the left femoral vein and of the inferior vena cava is essentially unchanged.

the renal veins are thrombosed. The contrast medium did not reach the height of the renal veins so I cannot tell whether there is washout by the blood coming from the renal veins to the vena cava nor was there any attempt to catheterize the renal

number of red blood cells was noted in the urine, a finding unusual in patients with membranous glomerular disease and perhaps related to anticoagulant therapy. A repeat renal biopsy was performed at this time and revealed changes suggestive of active glomerulonephritis. A transition of one type of renal lesion into another is very unusual in patients with SLE in terms of both our own experience⁴ and that of others.¹¹

Two years later the patient presented with another major catastrophe in acute myocardial infarction, at the remarkably young age of 21 years. A Type II hyperlipoproteinemia and persistent abnormalities of the nephrotic syndrome were found, but there were no immunologic signs of active disease. How can we explain the occurrence of acute myocardial infarction in this patient? SLE may produce a vasculitis involving the coronary arteries. However, large coronary artery occlusion with frank myocardial infarction is distinctly unusual.¹² The patient had hypercholesterolemia and Type II hyperlipoproteinemia. Serum lipid abnormalities are associated with the nephrotic syndrome regardless of etiology.⁷ Levels of cholesterol and triglyceride are increased due to enhanced hepatic synthesis of these lipids and retarded catabolism. Types II and IV lipoprotein abnormalities are most commonly found. There is an increased risk of atherosclerotic occlusive events in patients with the nephrotic syndrome. Of 36 patients with membranous glomerulonephritis at our own institution, four developed fatal myocardial infarction at a relatively young age.⁸ Berlyne and Mallick⁹ estimated that the risk of fatal infarction was increased 85 times in nephrotic patients.

Now we come to the last admission. He reentered the hospital in July 1971 with symptoms of congestive heart failure and testicular pain. Physical findings revealed a skin rash, more likely due to a superficial mycosis than to SLE, signs of congestive failure with an S_3 and S_4 gallop, and epididymitis. May we review the ECGs and x-rays at this time?

DR H. COHEN The first available ECG on this patient was on Nov 5 1969 (Fig 1). It is within normal limits. On April 9, 1971, slurred Q waves have developed in

Leads II, III, and V_6 . The QRS complex is now mainly inverted in Leads I and V_1 and biphasic in aV_L . ST segments are elevated in Leads II, III, and V_3 through V_6 and T waves are inverted in these leads. These changes are compatible with an acute posterolateral myocardial infarction probably with some loss of lateral myocardium as well, accounting for the slight right axis deviation. Almost four months later, on July 29, 1971, QRS complexes have become widened and slurred, especially in Leads I, V_1 , and V_6 . Q waves are no longer seen in Leads II, V_3 , and V_6 . The T waves are no longer inverted but ST segments remain elevated in Leads II, III, and the midprecordial leads. Thus an incomplete left bundle branch block has obscured part of the features of the myocardial infarction. In addition ST segment deviation has occurred partly because of the left bundle branch block, and partly because of digitalis effects, which may elevate ST segments in right precordial leads as well as in leads reflecting myocardial infarction such as Leads II and III in this case. Left bundle branch block disappears by the time of the next tracing (Aug 2 1971) and Q waves are again clearly seen in Lead II so that the inferior myocardial infarction is more obvious. In addition T waves have become inverted in the midprecordial leads. This suggests an active process in the inferior myocardium. This process may or may not be ischemic in nature. Also ectopic ventricular beats have developed. The last two tracings on Aug 11 and 18 1971 show a stable pattern of inferior myocardial infarction, digitalis effects, the development of left ventricular hypertrophy and widening and waving of anterior ischemic-like alterations. I feel that the deviation of the ST segments is still on the basis of digitalis effects and left ventricular hypertrophy if there is a pericarditis here. I am unable to recognize it as such mainly because of absence of the expected evolution and presence of other causes for the contour alterations.

DR SOMMERLERS Thank you Dr Cohen.

DR B. HAVIN First films of bony structure demonstrate what is noted in the protocol that is the steroid effects on the bones: demineralization and compression of the dorsal and lumbar vertebral bodies.

June 86
Number 4

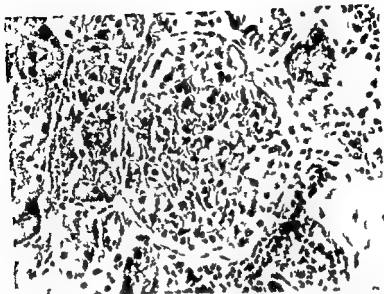


Fig 4 First renal biopsy (April 1965) Representative glomerulus showing diffuse membranous changes but no hypercellularity. Small interstitial infiltrates of round cells were also present (Hematoxylin and eosin $\times 490$)

suggest this possibility for several reasons. The description of the initial examination during his admission in July indicates that the heart tones were of good quality and no murmurs were heard. On the last admission he was admitted for congestive heart failure which was managed easily with the usual regimen of diuretics, digitals and salt restriction since he improved greatly within one day. On the second hospital day, however, he developed severe substernal chest pain with associated shortness of breath, diaphoresis and transient hypotension. The situation seemed to have been uncertain enough so that he was admitted to our coronary surveillance unit with the presumptive diagnosis of a possible acute myocardial infarction. He then developed a pericardial friction rub which certainly could be due to lupus pericarditis but is very commonly seen about a week after myocardial infarction in patients with myocardial infarction on an atherosclerotic basis alone. In addition his heart tones now were noted to be muffled whereas as I pointed out before on admission they were well heard. Furthermore he developed a holosystolic murmur that was not present on admission. Under these circumstances we usually ascribe the holosystolic murmur that results from infarction

or ischemic injury to either papillary muscle dysfunction or the development of a ventricular septal defect. The patient was discharged three weeks after this possible myocardial infarction. We do not know the level of his activity after discharge but the two to three week period after myocardial infarction if you will grant the possibility of an acute myocardial infarction is the time when we see myocardial ruptures or development of further infarction with arrhythmias and sudden death. So although I think your diagnosis is very well taken I would consider strongly the possibility of coronary atherosclerosis.

DR KANTLER: The protocol was somewhat unfair in that it failed to state that the patient had persistent angina pectoris between the second and final hospitalizations. He may not have reported this to the house staff. He was a very frightened young man who knew he was going to die. The diagnosis of his initial myocardial infarction was not easy. It was the fact that patients with long standing nephrotic syndrome are predisposed to arterio-sclerotic disease and myocardial infarction which influenced our admitting diagnosis. He presented with none of the classical features of infarction having developed abdominal distress during a fight with his mother and girl friend.

veins. Of interest, though, is that on first study I neglected to point out that we do see the calyceal system of both kidneys and they look intact. The kidneys however are somewhat enlarged, measuring 15 cm in length. One can see here the calyceal system on the left side and on the right side. They appear grossly normal except that the kidneys are large.

DR SIMON: Laboratory studies initially suggested inactive disease, but immunologic parameters of activity later became abnormal. The hospital course was characterized by recurrent episodes of shortness of breath, chest pain, transient hypotension, the appearance of a pericardial friction rub, and deterioration of renal function, suggesting a relentless downhill course. However his condition improved after treatment with digitalis, diuretics and heparin to the point where he was able to leave the hospital only to die suddenly two days later.

It appears that we must focus on the nature of the cardiac involvement in this patient with SLE. In view of the history of previous myocardial infarction, coronary atherosclerosis comes to mind first. But in spite of the stormy course in the hospital, serial ECGs and serum enzyme studies failed to establish the occurrence of fresh infarction. Therefore I doubt that coronary atherosclerosis could account for the sequence of events during the last hospital admission.

Pulmonary embolism is of course possible and was a source of serious concern to his physicians, especially in view of the history of thrombosis of the inferior vena cava and leg veins. Although it would be difficult to exclude pulmonary embolism, significant findings are not compatible with this diagnosis—the pericardial friction rub and negative lung scan. In one report on pulmonary embolism, positive lung scans were obtained in all patients.¹⁰

Finally, we may consider direct involvement of the heart by SLE. There may be endocarditis affecting the valve leaflets, papillary muscles, and mural endocardium, initially described by Libman and Sicks.¹¹ Although endocarditis may produce systolic and diastolic murmurs, it rarely is of sufficient hemodynamic consequence to result in congestive heart failure. In addi-

tion, endocarditis was not found in patients treated with corticosteroids.⁵

Myocardial damage may occur in SLE,¹² this is due to vasculitis of the small coronary arteries rather than to primary involvement of myocardial fibers. However, clinical evidence of frank myocardial infarction is unusual. The findings and course in the patient under discussion are entirely compatible with active lupus myocarditis.

Pericarditis is the most common manifestation of cardiac involvement in SLE,¹³ although not always evident clinically. The patient presented today did have a pericardial friction rub. Other alterations which contribute to cardiomegaly and congestive heart failure in patients with SLE are systemic hypertension, pulmonary hypertension from parenchymal or small vessel involvement in the pulmonary tree, anemia, and fever.

Therefore, I would conclude that the patient had systemic lupus erythematosus, active primarily involving the heart and the kidneys. The cardiac involvement was in the form of pericarditis and myocardial damage due to coronary vasculitis superimposed on atherosclerosis of the coronary arteries. Examination of the kidneys will show membranous glomerulonephropathy with an unusual transition to an active glomerulonephritis. As to the immediate cause of death, one can only speculate since the patient was not under observation that the mechanism was an arrhythmia or pulmonary embolism.

DR SOMMER: Dr Simon, that was the most lucid discussion that we have heard in many a year. I wonder if any of the people involved in the care of this patient before his demise would care to make a comment.

DR KANTER: I enjoyed your discussion very much. Dr Simon, I would like to know if you have any thoughts as to what we might have done during that last admission to determine the etiology of his cardiac disease.

DR SIMON: I will answer that after the pathologic report.

DR G. Glick: I also enjoyed very much your discussion, Dr Simon. Just on the basis of this protocol, I would think that coronary atherosclerosis may be the primary reason for the patient's death. I



Fig 7 Second renal biopsy Electron micrograph of a glomerular capillary There were extensive electron-dense subepithelial deposits (arrows) Smaller deposits were also present within the basement membrane proper (BV) and in the mesangial areas (M) The capillary lumen (L) is narrowed by the swollen endothelial cells (EN) (X8400)

First I will present the findings of the two renal biopsies One was obtained in 1965 at the University of Illinois It was an adequate biopsy and all glomeruli showed essentially the same changes namely a diffuse thickening of the capillary wall with normal cellularity and patent capillary lumens (Fig 4) In periodic acid-Schiff and better in silver methenamine preparations a true thickening of the basement membrane and some irregularity of its contour could be seen (Fig 5) There was a mild increase of mesangial matrix but this was not thought to be significant Under oil immersion it was possible to recognize projections or spikes from the outer aspect of the basement membrane In trichrome preparations under oil immersion on the basement membrane and its projection from the outer aspect were blue Separating the spikes were fuchsin-

philic red areas corresponding to immune deposits In brief the histologic pattern of the first biopsy was quite consistent with either that of idiopathic membranous glomerulopathy or that of the membranous form of lupus nephritis^{1,2}

The second renal biopsy was obtained about four years later The disease appeared to be more active The glomerular changes were similar to those of the first biopsy except for an occasional small epithelial crescent (Fig 6) There was now an interstitial infiltrate of inflammatory cells for the most part lymphocytes and plasma cells which should be interpreted as a sign of activity of lupus nephritis There was more interstitial edema and fibrosis as well and in the glomerular capillaries few trapped leukocytes could be seen We must keep in mind that at this time the patient had also a reasonably



Fig 5 First renal biopsy. The glomerular capillary basement membrane is diffusely thickened and its outer contour is irregular (Silver methenamine $\times 1235$)

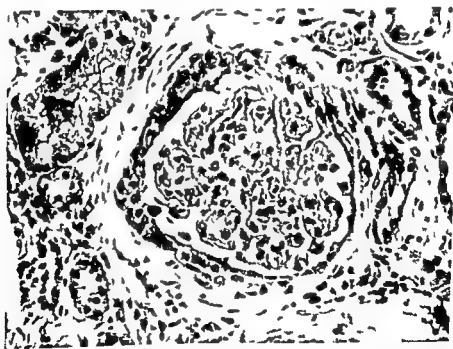


Fig 6 Second renal biopsy (November 1969). All glomeruli showed membranous changes as in the first biopsy but in addition some of them presented small epithelial crescents as shown in this figure. There was also some interstitial edema and tubular atrophy (Hematoxylin and eosin $\times 490$)

He was admitted to the hospital despite the protestations of the house staff who attributed his complaints to anxiety following a family argument. Of course the ECG settled the matter.

DR C I PIRANI This case was selected for a CPC first because of its long and in-

teresting history and second because all the major clinical manifestations of this patient's illness were satisfactorily explained by the postmortem examination. Dr Simon has given us a very lucid presentation nicely correlated with possible pathologic findings and this will make my task much easier.

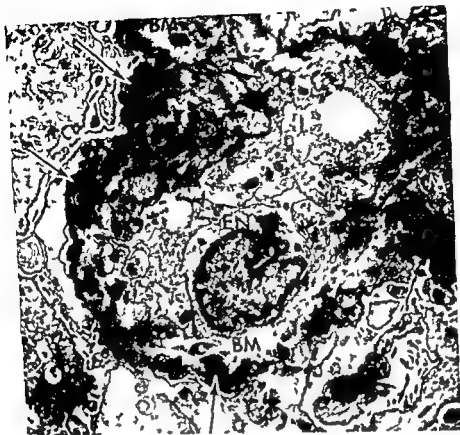


Fig 7 Second renal biopsy Electron micrograph of a glomerular capillary There were extensive electron dense subepithelial deposits (arrows) Smaller deposits were also present within the basement membrane proper (BM) and in the mesangial areas (M) The capillary lumen (L) is narrowed by the swollen endothelial cells (E) ($\times 8400$)

First I will present the findings of the two renal biopsies. One was obtained in 1965 at the University of Illinois. It was an adequate biopsy and all glomeruli showed essentially the same changes, namely, a diffuse thickening of the capillary wall with normal cellularity and patent capillary lumens (Fig 4). In periodic acid-Schiff and better in silver methenamine preparations, a true thickening of the basement membrane and some irregularity of its contour could be seen (Fig 5). There was a mild increase of mesangial matrix, but this was not thought to be significant. Under oil immersion it was possible to recognize projections or spikes from the outer aspect of the basement membrane. In trichrome preparations under oil immersion the basement membrane and its projections from the outer aspect were blue. Separating the spikes were fuchsin-

philic red areas corresponding to immune deposits. In brief, the histologic pattern of the first biopsy was quite consistent with either that of idiopathic membranous glomerulopathy or that of the membranous form of lupus nephritis.¹²

The second renal biopsy was obtained about four years later. The disease appeared to be more active. The glomerular changes were similar to those of the first biopsy except for an occasional small epithelial crescent (Fig 6). There was now an interstitial infiltrate of inflammatory cells, for the most part lymphocytes and plasma cells, which should be interpreted as a sign of activity of lupus nephritis. There was more interstitial edema and fibrosis as well, and in the glomerular capillaries few trapped leukocytes could be seen. We must keep in mind that at this time the patient had also a reasonably

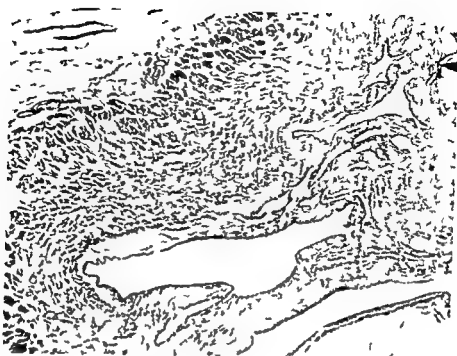


Fig 8 Cross section through the right renal vein near the inferior vena cava. A well organized and recanalized thrombus is present (Hematoxylin and eosin $\times 40$).

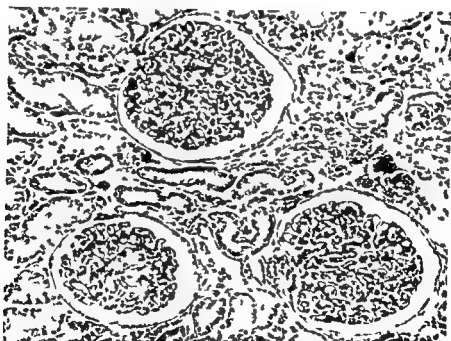


Fig 9 At autopsy, moderate membranous changes were present in all glomeruli. No changes of note were seen either in the tubules or in the interstitium (Hematoxylin and eosin $\times 200$).

well documented episode of renal vein thrombosis which may have been responsible for the prominent interstitial edema.¹⁴ Electron microscopy of the second biopsy revealed the capillaries of all glomeruli studied to have a very thick basement membrane with electron dense deposits primarily in a subepithelial position but

also within the basement membrane proper. Electron dense deposits were also seen in the mesangial areas.¹⁵ The visceral epithelial cells showed villous transformation and the foot processes were generally fused (Fig. 7). The endothelial cells were often swollen and occasionally were found to contain viruslike inclusions in their cyto-

plasm—a common finding in SLE.¹⁶ In brief the electron microscopic studies confirmed the diagnosis of lupus nephritis membranous type.

At the time of autopsy the external examination revealed a maculopapular skin lesion which histologically was found to be due to a mycotic infection probably trichophytosis and which we thought to be related to the low immunologic resistance of this patient. There was a moderate amount of aortic atherosclerosis especially in the abdominal segment. The inferior vena cava below the renal veins presented an irregular intimal surface due to a web of fibrous tissue. The right renal vein also contained some bands of fibrous tissue. However the lumen of both these vessels was adequately patent. Histologically these bands were obviously part of a recanalized thrombus (Fig. 8). The kidneys were large and had a smooth surface. Histologically the renal changes were quite similar to those observed on the second renal biopsy except that all evidence of active disease was no longer present. Actually the tubules were well preserved and there was little interstitial edema or fibrosis (Fig. 9). There was however a mild to moderate degree of nephrocalcinosis in the renal medulla. This almost certainly was related to corticosteroid therapy as was the extremely severe osteoporosis seen in sections from the vertebrae confirming the x-ray findings. As we have occasionally seen in patients with osteoporosis even with normal serum calcium levels there was a mild degree of parathyroid hyperplasia. However no changes of osteitis fibrosa or of increased osteoclastic activity were seen. The spleen showed atrophic lymphoid tissue and the adrenal cortices were somewhat reduced in size both changes obviously secondary to corticosteroid therapy. The heart weighed 410 grams much of this increased weight due to the abundant subepicardial fat. Thus hypertrophy was considered to be only mild to moderate. On cross section the right coronary artery a few centimeters from its orifice was seen to be occluded by a recent thrombus. Histologically the thrombus could be seen to have formed over an atheromatous plaque (Fig. 10). The left circumflex coronary artery was completely occluded by an old athero-



Fig. 10 Cross section through the right coronary artery showing extensive atheromatous changes and a superimposed recent thrombus (Hematoxylin and eosin $\times 65$)

sclerotic process with calcification. Other coronary arteries presented considerable atherosclerosis. There was a rather large infarct in part old and in part recent involving primarily the posterior aspects of the interventricular septum and of the left ventricular wall. The left papillary muscles all the way to their apex were involved by fibrosis and by very recent ischemic necrosis probably no more than two days old (Fig. 11). This lesion was almost certainly responsible for the holosystolic murmur heard a few days before death. A small organizing mural thrombus was attached to the area of infarction in the left posterior wall which was estimated to be six to eight weeks old. To complicate matters in the pericardial adipose tissue a few small arteries presented the classic changes of acute arteritis. A possibility that Dr. Simon mentioned. The inflammatory cells consisting of polymorphonuclear leukocytes, lymphocytes and plasma cells extended through the entire thickness of the wall in which fibroid changes were also present (Fig. 12). No microorganisms were seen with the use of special stains. The possi-

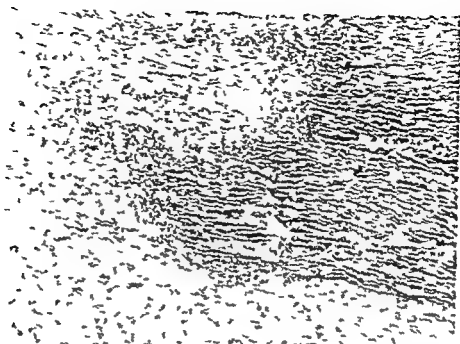


Fig. 11 Section through a left papillary muscle showing recent myocardial necrosis (right) and an organizing infarct (left) (Hematoxylin and eosin $\times 52$)

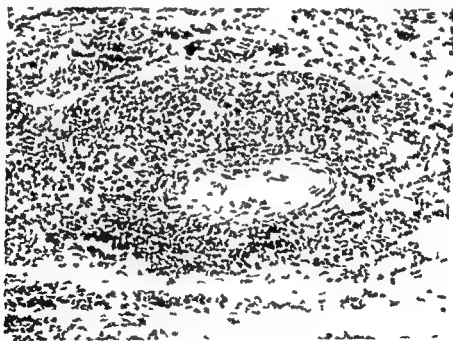


Fig. 12 Peridrenal arteriole with acute inflammatory changes involving its wall (Hematoxylin and eosin $\times 220$)

bility of hypersensitivity was considered but no eosinophils were present in the inflammatory infiltrates and no arteritic lesions were seen in any other tissue. Death was attributed to heart failure as evidenced by the severe edema and congestion of the lungs. So much of the symptomatology of this patient can be satisfactorily explained by the autopsy findings.

This patient for 19 years or more had a form of lupus nephritis which has a very good prognosis indeed not dissimilar from that of idiopathic membranous glomerulonephropathy. These patients can live a long life as long as the other symptomatology and complications of SLE can be controlled. It is of interest that the episode of renal vein thrombosis appeared to have

aggravated the renal disease only temporarily. This may have been due to the prompt diagnosis and treatment of this complication. As a result of the nephrotic syndrome and persistent hyperlipidemia, severe atherosclerosis, particularly of the coronary arteries, developed early in the life of this patient and led to the occlusion of two major coronary arteries and extensive myocardial infarction. The possibility that periarteritis may have facilitated the development of atherosclerosis does not seem likely. The isolated periaortic arteritis remains unexplained but there was no evidence of either old or recent arteritis in the coronary arteries.

DR SIMON: I steered away from the diagnosis of coronary atherosclerosis as the primary cardiac lesion because of the finding of a pericardial friction rub and the absence of confirmatory evidence of myocardial infarction. This line of thinking was based on the assumption that the presence of a pericardial friction rub in a patient with myocardial infarction would indicate a transmural infarct which should be reflected in diagnostic ECG and serum enzyme changes. It is sobering to realize that our constructive ways of thinking are not always valid and that Dr Glick's summation was more accurate than my own.

An important issue raised by this case is the need to alter our approach to the patient with the nephrotic syndrome in that attention must be paid not only to the kidneys but also to the significant lipid and coagulation abnormalities that are present with the enhanced risk of atherogenesis and thrombotic events. If treatment of the underlying renal lesion produces a remission of proteinuria, we may expect resolution of the associated lipid and clotting abnormalities. However, if treatment is unsuccessful in altering proteinuria, then direct therapeutic efforts at lowering serum lipids by diet and drugs and preventing thrombosis through the use of anticoagulants should be strongly considered.

In a recent report, arteritis of a large coronary artery with occlusion and myocardial infarction was demonstrated in a 16-year-old girl with SLE.¹² In addition, angiographic abnormalities of major coronary vessels were found in three other

young women. All had angina pectoris and two sustained myocardial infarctions.

DR S. KABINS: I just want to ask Dr Glick whether somewhere in the last month or so before he died, this patient might have been a candidate for coronary angiography or coronary bypass surgery?

DR GLICK: The answer to both questions is yes. I would have considered him a candidate for both angiography and surgery. However, I have great doubt that either of these procedures would have been carried out. We do coronary angiography only when we think corrective surgery is in the cards and in view of his primary disease, I doubt that we would have thought that surgery would have been helpful.

DR SOMMERS: Any more comments or questions from the audience? If not, thank you very much.

REFERENCES

1. Simon N M, Werner W J, Franklin W A, and Potter E V: Comparison of steroid and combined steroid immunosuppressive treatment of lupus erythematosus nephritis. *Ann Intern Med* 74:847, 1971.
2. Pollak V E and Pirani C L: Renal histologic findings in systemic lupus erythematosus. *Mayo Clin Proc* 44:630, 1969.
3. Baldwin D S, Lowenstein J, Rothfield N F, Gallo G, and McCluskey M T: Clinical course of proliferative and membranous forms of lupus nephritis. *Ann Intern Med* 73:929, 1970.
4. Kendall A G, Lehmann R C, and Dossetor J B: Nephrotic syndrome: a hypercoagulable state. *Arch Intern Med* 127:1021, 1971.
5. Shearn M A: The heart in systemic lupus erythematosus. *AM HEART J* 58:452, 1959.
6. Bridgman W, Bywaters F G L, Lessof M H, and Ross I P: The heart in systemic lupus erythematosus. *Br Heart J* 22:1, 1960.
7. Simon N M and del Greco F: Lipid abnormalities in renal disease. *Ann Clin Lab Sci* 2:186, 1972.
8. Earle D P: Glomerulonephritis: clinical aspects. *Bull N Y Acad Med* 46:749, 1970.
9. Berlyne G M and Mallick N P: Ischemic heart-disease as a complication of nephrotic syndrome. *Lancet* 2:399, 1969.
10. Szucs M M, Jr, Brooks H L, Grossman W, Bana J S, Jr, Meister S G, Dexter L, and Dalen J W: Diagnostic sensitivity of laboratory findings in acute pulmonary embolism. *Ann Intern Med* 74:161, 1971.
11. Libman E and Sacks M A: A hitherto undescribed form of valvular and mural endocarditis. *Arch Intern Med* 33:701, 1924.
12. Bonfio T A, Botti R E, and Hagstrom J W C: Coronary arteritis: occlusion and

Table 1 *Classification of cases of CAVO without splenic abnormalities and their relationship to Down's syndrome*

| Complete | | | | | Various complexes of CAVO | Incomplete | | | | |
|---|----|----|----|-------|---|------------|----|---|---|---|
| N† | D† | U† | T† | Total | | Total | V | D | U | T |
| A In levocardia (normal position of the heart) with situs solitus organs | | | | | | | | | | |
| 1 Simple (without other complexes) | | | | | | | | | | |
| 15 | 30 | 2 | 0 | 47 | a Balanced | 11 | 9 | 2 | 0 | 0 |
| 2 | 1 | 0 | 0 | 3 | b Dominant right | 3 | 3 | 0 | 0 | 0 |
| 3 | 1 | 0 | 1 | 5 | c Dominant left | 1 | 1 | 0 | 0 | 0 |
| 55 | | | | | | 15 | | | | |
| 2 Complicated (associated with other complexes) | | | | | | | | | | |
| 7 | 7 | 0 | 0 | 14 | a Left coarctation | 10* | 10 | 0 | 0 | 0 |
| 0 | 8 | 0 | 0 | 8 | b Patent ductus arteriosus | 1 | 0 | 1 | 0 | 0 |
| 1 | 0 | 0 | 1 | 2 | c Pulmonary stenosis | | | | | |
| 4 | 10 | 1 | 0 | 15 | d Tetralogy of Fallot | 1 | 0 | 1 | 0 | 0 |
| 7 | 0 | 1 | 0 | 8 | e Double outlet right ventricle | 1 | 1 | 0 | 0 | 0 |
| 3 | 0 | 1 | 0 | 4 | f Complete transposition | | | | | |
| 9 | 3 | 1 | 0 | 13 | g Absence of atrial septum | | | | | |
| 0 | 1 | 0 | 0 | 1 | h Absence of posterior ventricular septum | | | | | |
| | | | | | i Aortic stenosis | 1 | 1 | 0 | 0 | 0 |
| 4 | 0 | 0 | 0 | 4 | j Total or partial anomalous pulmonary venous drainage | 1 | 0 | 1 | 0 | 0 |
| (partial) | | | | | k Single ventricle | (total) | | | | |
| 2 | 0 | 5 | 0 | 7 | B In mixed dextrocardia (atria pivotal ventricles in mirror image) | | | | | |
| 1 | 0 | 0 | 0 | 1 | C In isolated levocardia with atrial situs inversus | | | | | |
| 2 | 0 | 0 | 0 | 2 | D In isolated levocardia with atrial undeterminate with situs inversus organs | | | | | |
| 2 | 0 | 0 | 0 | 2 | L In mesocardia with atrial situs solitus and ventricles in mesoversion | | | | | |
| 1 | 0 | 0 | 0 | 1 | I In mesocardia—indeterminate type | | | | | |
| 0 | 1 | 0 | 0 | 1 | | | | | | |
| 83 | | | | | | 15 | | | | |

*Includes 2 cases of adult coarctation

†Abbreviations: D = Down's syndrome; N = Down's syndrome absent; T = Trisomy 21; U = Unknown whether Down's syndrome present or not

dominant left form (Fig. 2), the left ventricle is markedly hypertrophied and enlarged, but the right ventricle is smaller than normal, with a normal or thickened wall. These three forms were found in both types complete and incomplete and most complexes of CAVO. In the incomplete type in the complexes with fetal coarctation, and in all types of dextrocardia, there was a greater tendency for the occurrence of the right ventricular form than in the other complexes.

The term *mixed dextrocardia* (in Tables I, II, and III) implies that the base apex of the heart points to the right and caudal ward and the atria and ventricles do not

correspond. The term *presumptive* in *Mirror image dextrocardia* (Table II) implies that the atrial septum is absent or inconclusive in its morphology as to its right or left side and hence only a presumptive diagnosis is made of the atria from the atrial appendages.

In consideration of abnormalities of the spleen all abnormalities were included in this study both minor and major. Even though accessory spleens and occasionally abnormal lobulation of the spleen may occur in the normal person yet they may conceivably represent a forme fruste of asplenia and polysplenia, and hence they are included in this study.

Table 11 Classification of cases of CAVO with aplenic abnormalities and their relationship to Down's syndrome

| Complete | | | | | | | | | | | | Incomplete | | | | | |
|---|---|---|---|---|---|----|---|---|---|----|-------|---------------------------|--|--|--|--|--|
| N | D | U | T | A | P | AC | B | T | H | MA | Total | Various complexes of CIVO | | | | | |
| N | D | U | T | A | P | AC | B | T | H | MA | | | | | | | |
| A In levocardia with situs solitus organs | | | | | | | | | | | | | | | | | |
| 1 Simple | | | | | | | | | | | | | | | | | |
| 2 Balanced | | | | | | | | | | | | | | | | | |
| 3 Complicated | | | | | | | | | | | | | | | | | |
| a Patent ductus arteriosus | | | | | | | | | | | | | | | | | |
| b Patent ductus arteriosus | | | | | | | | | | | | | | | | | |
| c Pulmonary stenosis | | | | | | | | | | | | | | | | | |
| d Double outlet right ventricle | | | | | | | | | | | | | | | | | |
| e Complete transposition | | | | | | | | | | | | | | | | | |
| f Aortic atresia and interrupted aortic arch | | | | | | | | | | | | | | | | | |
| g Single ventricle | | | | | | | | | | | | | | | | | |
| h Truncus | | | | | | | | | | | | | | | | | |
| B In dextrocardia | | | | | | | | | | | | | | | | | |
| 1 Presumptive mirror image type | | | | | | | | | | | | | | | | | |
| 2 Mixed dextrocardia (atrial pivotal ventricles in mirror image) | | | | | | | | | | | | | | | | | |
| 3 Type undetermined | | | | | | | | | | | | | | | | | |
| C In isolated levocardia | | | | | | | | | | | | | | | | | |
| 1 With situs solitus | | | | | | | | | | | | | | | | | |
| 2 With indeterminate atria | | | | | | | | | | | | | | | | | |
| D In meocardia | | | | | | | | | | | | | | | | | |
| 1 Presumptive mesoversion with atrial situs solitus and ventricles in mesoversion | | | | | | | | | | | | | | | | | |
| 2 Mixed meocardia with atrial situs solitus and ventricles inverted | | | | | | | | | | | | | | | | | |
| 3 Indeterminate type | | | | | | | | | | | | | | | | | |
| 7 | | | | | | | | | | | | | | | | | |

Table III *Classification of cases of CAVO in which the status of the spleen is unknown* and their relationship to Down's syndrome*

| Complete | | | | | Various complexes of CAVO | Incomplete | | | | |
|---|----|----|----|-------|---|------------|---|---|---|---|
| A† | D† | U† | T† | Total | | Total | N | D | U | T |
| | | | | | | | | | | |
| A In levocardia with situs solitus organs | | | | | | | | | | |
| 1 Simple | | | | | | | | | | |
| 3 | 3 | 7 | 0 | 13 | a Balanced | 6 | 5 | 0 | 1 | 0 |
| 0 | 1 | 0 | 0 | 1 | b Dominant right | | | | | |
| | | | | | c Dominant left | | | | | |
| 2 Complicated | | | | | | | | | | |
| 0 | 1 | 0 | 0 | 1 | a Fetal coarctation | 1 | 1 | 0 | 0 | 0 |
| 1 | 2 | 0 | 0 | 3 | b Patent ductus arteriosus | | | | | |
| 1 | 0 | 0 | 0 | 1 | c Pulmonary stenosis | | | | | |
| 1 | 0 | 2 | 0 | 3 | d Tetralogy of Fallot | | | | | |
| 0 | 0 | 1 | 0 | 1 | e Double outlet right ventricle | | | | | |
| 0 | 0 | 1 | 0 | 1 | f Complete transposition | | | | | |
| 2 | 0 | 1 | 0 | 3 | g Absent atrial septum | | | | | |
| 0 | 0 | 1 | 0 | 1 | h Total anomalous pulmonary venous drainage | | | | | |
| 0 | 0 | 1 | 0 | 1 | i Truncus | | | | | |
| 1 | 0 | 2 | 0 | 3 | j Single ventricle | | | | | |
| B In dextrocardia | | | | | | | | | | |
| 0 | 0 | 2 | 0 | 2 | Mixed dextrocardia with atria situs solitus and ventricles inverted | | | | | |
| — 34 | | | | | | — 7 | | | | |

*In this group the status of the spleen could not be ascertained because (1) only a chest autopsy was done or (2) the autopsy protocol did not distinctly indicate what the status of the spleen was or (3) the autopsy protocol was not available.

†Abbreviations: D = Down's syndrome; N = Down's syndrome absent; T = Trisomy 18; U = Unknown whether Down's syndrome present or not.

Table IV *Occurrence of Down's syndrome in the various complexes of CAVO**

| Type | Occurrence |
|---|------------|
| Complete simple without splenic abnormalities | 60.4% |
| Complete complicated without splenic abnormalities | 40.5% |
| Complete with splenic abnormalities | 7.3% |
| Incomplete simple without splenic abnormalities | 13.3% |
| Incomplete complicated without splenic abnormalities | 20.0% |
| Incomplete with splenic abnormalities | 28.6% |
| Complete with positional abnormalities of the heart and organs with normal spleen | 14.3% |
| Complete and incomplete with positional and splenic abnormalities | 0.0% |

*Percentages do not indicate incidence of the anomalies in Down's syndrome.

The ductus arteriosus is mentioned as being patent only if it is so after 3 months of age. Below 3 months of age it is considered to be normal.

The AV valve in CAVO

The three types of anterior bridging leaflets seen in the complete form de-

scribed by Rastelli and colleagues^{28, 35, 41} were also found in our material. In their first type there are attachments of the anterior bridging leaflet by chordae to both sides of the ventricular septum. In their second type the anterior bridging leaflet was unattached to the septum but attached medially to an anomalous papillary muscle in

6.1 me 86
1 mb 4



Fig 1 A and B Dominant right form of CAVO intermediate type A Right atrial and right ventricular view B Left atrial and left ventricular view RA = right atrium RV = right ventricle LA = left atrium LV = left ventricle



Fig 2 A and B Dominant left form of CAVO complete type A Left atrial and left ventricular view B Right ventricular view LA = left atrium LV = left ventricle A = aorta RV = right ventricle PA = pulmonary trunk

the right ventricle. In their third type the anterior bridging leaflet was a free floater being unattached to the septum above but attached to the usual papillary muscles on both sides. The first and third types of these authors however are not sharply demarcated as categories in our material. Some free floating leaflets were completely free floating, with connections of the leaflet only to the left anterior and right anterolateral papillary muscles with no

muscle of Lancisi being present. Others had a few chordal attachments to the summit of the ventricular septum either on the right or left side. Still others had wide attachments of chordae to both sides of the summit. The annulus of this leaflet likewise had a varying morphology. It was usually attached to the non-coronary and to the adjacent part of the left coronary cusp by way of the *pars membranacea* and what may be called the aortico-mitral annulus. The

connection in some cases involved also the posterior extremity of the right aortic cusp and in a rare case extended throughout the whole base of the right aortic cusp with a varying amount of attachment on the summit of the anterior basal portion of the ventricular septum. The morphology of the posterior bridging and lateral leaflets was as described by Rastelli and co workers.^{13,25,43}

The right anterolateral papillary muscle was usually widely attached to the wall often had multiple components, and in some cases was prolonged extending into the annulus of the anterior bridging leaflet.

The right inferior papillary muscle in some cases was absent while in others there were several such structures. The left anterior and posterior papillary muscles were in some cases situated more laterally than normal. These muscles were in some instances small and occasionally they proceeded to the base of the valve rather than to the edge. Only in occasional case of the second type of Rastelli and co workers^{25,35,43} was seen with a small papillary muscle attachment to the right side of the septum. Two cases of double orifices were seen on the mitral side and 3 cases of double orifices on the tricuspid side.

In the incomplete (intermediate transitional) type as described by Wikström and Edwards,⁶ valvular tissue was seen on the summit of the ventricular septum. In some cases this was an extension of the posterior bridging leaflet. In others it appeared to be an independent component.

Alterations in the annular relationships of the anterior bridging leaflet were found in CAVO associated with tetralogy of Fallot, double outlet right ventricle and in complete transposition. In tetralogy of Fallot the anterior bridging leaflet was almost always a free floater. In some of these cases the base of the leaflet was related more to the left than to the non-coronary cusp. The right anterolateral papillary muscle was usually abnormally large, widely based, and its chordae often formed a sheet continuing muscle extending to the base of the valve. In double outlet right ventricle and in partial transposition with pulmonary atresia (*pseudotruncus*) the anterior bridging leaflet was always a free floater.¹ Its annulus was either related to the aorta by way of the *pars membranacea*

or to an extension of the central fibrous body, or it was separated from both the aorta and the pulmonary trunk by conal musculature. In complete transposition there was almost always pulmonary atresia and the base of the anterior bridging leaflet in almost all cases was separated from the aorta by conal musculature. In complicated types in general there was a strong tendency for the anterior bridging leaflet to be a "free floater."

Alterations of the attachments of the common AV valve to the papillary muscles were found in single ventricle. Here there was usually a group of anterior and posterior papillary muscles with a varying number of leaflets.

The attachment of the leaflets in the dominant right and left forms deserves attention. In the latter in many cases the only attachment in the right ventricle was that of the anterior bridging leaflet to the anterolateral papillary muscle while the remainder of the valve was attached to the left ventricle. In the dominant right type, the attachment of the leaflets were in the balanced form but the left portions were small and the right large.

Discussion

In our material without splenic abnormality CAVO both complete and incomplete types occurs as a complex by itself or associated with other complexes among which most common are fetal connection patent ductus arteriosus tetralogy of Fallot double outlet right ventricle absence of the atrial septum and single ventricle. Less commonly it is associated with pulmonary stenosis absence of the posterior part of the ventricular septum complete transposition aortic stenosis or atresia interruption of the aortic arch and truncus communis. It may be noted that despite a normal spleen it occasionally is associated with dextrocardia isolated levocardia and mesocardia. It is also of interest that in our material without splenic abnormalities, there are more cases of the complete complicated type than the complete simple type. If this should prove to be true for other series this needs to be kept in mind in the efficacy of surgical treatment.

In all types complete and incomplete simple or complicated with or without

asplenia there may be dominant right and dominant left forms. We have chosen to delineate the dominant right and dominant left forms of CAVO as those in which the smaller of the two ventricles is smaller than normal rather than those in which one chamber is larger than the other. We believe this is justified from the surgical standpoint. As long as a chamber is normal in size and thickness it is presumed that at least theoretically a case of CAVO may be operable. When however a chamber is smaller than normal there is some question as to the operability of the case.

The question of the tendency towards dominance of the right or the left sides in some cases of CAVO has not been explained. In favor of an embryological explanation for the dominant left form are the tenuous attachments of the valve to the right ventricle reminiscent of a primitive ventricle perhaps of the same type as found in the anomaly straddling tricuspid orifice.¹⁰ However there may be physiologic reasons for dominance of either side which have not yet come to the fore in correlative catheterization and anatomic studies.

The association of CAVO with splenic abnormalities is well known.¹¹ Our material shows that this may occur with normal or abnormal situs of organs. Where there is situs solitus in the complete type only rarely is the association found in the simple type or with fetal (preductal) coarctation and patent ductus arteriosus when the association is present there is usually only an accessory spleen or abnormal lobation of the spleen. However when the complete form in situs solitus is complicated by pulmonary stenosis double outlet right ventricle aortic atresia absence of the atrial septum or single ventricle the association with splenic abnormalities is frequent and there usually is asplenia. Hence in situs solitus the simple type of CAVO does not show severe abnormalities of the spleen but the complicated form may show such abnormality. The incomplete type simple or complicated in situs solitus is rarely associated with splenic abnormalities and then only with an accessory or multilobed spleen. Where there is abnormal situs of organs it is well known^{12,13} the association of CAVO with asplenia or polyplenia is strong.

As concerns the relationship of CAVO to Down's syndrome as a well known the complete simple type without splenic abnormalities shows a frequent association. As compared with the complete simple type this association is less marked in the complicated type ($p < 0.5$). As compared to the complete simple form there is a less frequent association of the incomplete type without splenic abnormalities with Down's syndrome ($p < 0.1$). Interesting is the relative lack of association of the complete type with Down's syndrome where there are abnormalities of the spleen as compared to CAVO without the latter ($P < 0.01$). The reason for this inverse relationship is unknown to us.

From the standpoint of longevity as gleaned from autopsy statistics the incomplete simple type has a greater longevity than the complete simple type. Fetal coarctation in CAVO is relatively lethal in both complete and incomplete types whereas the presence of a ductus arteriosus in the complete simple type may be beneficial. Splenic abnormalities in CAVO are associated with less viability than those cases with normal spleens.

Summary

A study was made of the incidence of CAVO as it occurs by itself and as it is associated with other complexes and the association of these complexes with Down's syndrome and splenic abnormalities. It was found that the complete complicated type (associated with other abnormalities) is more common than the complete simple type where there are no splenic abnormalities. All types complete and incomplete had balanced dominant right and dominant left forms which is important surgically. The simple type in situs solitus does not show severe abnormalities of the spleen but the complicated type does. In complete types are rarely associated with severe abnormalities of the spleen.

The complete complicated type without splenic abnormalities is less often associated with Down's syndrome than is the complete simple and the incomplete less often than the complete type. There is an inverse relationship between Down's syndrome and CAVO associated with splenic abnormalities.

connection in some cases involved also the posterior extremity of the right aortic cusp and in a rare case extended throughout the whole base of the right aortic cusp with a varying amount of attachment on the summit of the anterior basal portion of the ventricular septum. The morphology of the posterior bridging and lateral leaflets was as described by Rastelli and co workers.^{20, 25, 43}

The right anterolateral papillary muscle was usually widely attached to the wall often had multiple components and in some cases was prolonged extending into the annulus of the anterior bridging leaflet. The right inferior papillary muscle in some cases was absent while in others there were several such structures. The left anterior and posterior papillary muscles were in some cases situated more laterally than normal. These muscles were in some instances small and occasionally they proceeded to the base of the valve rather than to the edge. Only in occasional case of the second type of Rastelli and co workers^{20, 25, 43} was seen with a small papillary muscle attachment to the right side of the septum. Two cases of double orifices were seen on the mitral side and 3 cases of double orifices on the tricuspid side.

In the incomplete (intermediate transitional) type as described by Wiklund and Edwards,⁶ valvular tissue was seen on the summit of the ventricular septum. In some cases this was in extension of the posterior bridging leaflet. In others it appeared to be an independent component.

Alterations in the annular relationships of the anterior bridging leaflet were found in CAVO associated with tetralogy of Fallot, double outlet right ventricle and in complete transposition. In tetralogy of Fallot, the anterior bridging leaflet was almost always a 'free floater'. In some of these cases the base of the leaflet was related more to the left than to the non-coronary cusp. The right anterolateral papillary muscle was usually abnormally large, widely based and its chordae often formed a sheet containing muscle extending to the base of the valve. In double outlet right ventricle and in partial transposition with pulmonary atresia (*pseudotruncus*) the anterior bridging leaflet was always a 'free floater'. Its annulus was either related to the aorta by way of the *pars membranacea*

or to in extension of the central fibrous body, or it was separated from both the aorta and the pulmonary trunk by conal musculature. In complete transposition there was almost always pulmonary atresia and the base of the anterior bridging leaflet in almost all cases was separated from the aorta by conal musculature. In complicated types in general there was a strong tendency for the anterior bridging leaflet to be a 'free floater'.

Alterations of the attachments of the common AV valve to the papillary muscles were found in single ventricle. Here there was usually a group of anterior and posterior papillary muscles with a varying number of leaflets.

The attachment of the leaflets in the dominant right and left forms deserves attention. In the latter in many cases the only attachment in the right ventricle was that of the anterior bridging leaflet to the anterolateral papillary muscle, while the remainder of the valve was attached to the left ventricle. In the dominant right type the attachment of the leaflets were as in the bifurcated form but the left portions were small and the right large.

Discussion

In our material without splenic abnormality CAVO both complete and incomplete types occurs as a complex by itself or associated with other complexes among which most common are fetal correction, patent ductus arteriosus, tetralogy of Fallot, double outlet right ventricle, absence of the atrial septum and single ventricle. Less commonly it is associated with pulmonary stenosis, absence of the posterior part of the ventricular septum, complete transposition, aortic stenosis or atresia, interruption of the aortic arch and truncus communis. It may be noted that despite a normal spleen it occasionally is associated with dextrocardia, isolated levocardia and mesocardia. It is also of interest that in our material without splenic abnormalities there are more cases of the complete complicated type than the complete simple type. If this should prove to be true for other series this needs to be kept in mind in the efficacy of surgical treatment.

In all types complete and incomplete simple or complicated with or without

- repair of endocardial cushion defects Ann Thorac Surg 2:399 1966
- 31 Kurlander G L and Garod D A Oblique (left ventriculography) in endocardial cushion defect Grade III Am J Roentgenol Radium Ther Nucl Med 9 314 1964
- 32 Somerville J Clinical assessment of the function of the mitral valve in atrioventricular defects related to the anatomy AM HEART J 71:101 1966
- 33 Rastelli G C Kirklin J W and Kincaid O W Angiocardiography of persistent common atrioventricular canal Mayo Clin Proc 42:200 1967
- 34 Gerbode F Sanchez P A Aguero R Keith W J Hill D DeVries P A Selzer A and Robinson S Endocardial cushion defects Ann Surg 166:489 1967
- 35 Rastelli G C Ongley P A Kirklin J W and McGoon D C Surgical repair of the complete form of persistent common atrioventricular canal J Thorac Cardiovasc Surg 50:799 1968
- 36 Taguchi K Suaki N Okamoto Matsuura Y and Hrao M Surgical experience with persistent common atrioventricular canal in a series of eighty two patients particular consideration on the connection of valve incompetence and deficiency of ventricular septum J Thorac Cardiovasc Surg 50:501 1968
- 37 Baron M G Endocardial cushion defects Radiol Clin North Am 6:343 1968
- 38 Goor D Lillehei W and Edwards J E Further observations on the pathology of the atrioventricular canal malformations Arch Surg 97:954 1968
- 39 Emanuel R Nchois J Aaders J M Morris F C and Somerville J AV defects A study of 92 families Br Heart J 30:645 1968
- 40 d Allaines C Colvez P Levasseur P Facquet J and Dubost C A rare congenital cardiopathy association of tetralogy with complete AV communication Clinical detection and surgical repair Arch Mal Coeur 62:996 1969
- 41 Shih C Y Patel M K and Hastreter A R Hemodynamics of complete atrioventricular canal and its evolution with age Am J Cardiol 24:326 1969
- 42 Gilbert G Taillon J P Aérichide N and David P A study of 38 cases of atrioventricular canal Union Med Can 129:1 1968
- 43 Rastelli G C Ongley P A and McGoon D C Surgical repair of complete atrioventricular canal with anterior common leaflet undivided and unattached to ventricular septum Mayo Clin Proc 44:335 1969
- 44 Di Matteo J Galey J J Ribierre M Vacheron A Vanetti A Sabaut D and Lafont H Correction of a complete form of common atrioventricular canal Arch Mal Coeur 63:1042 1970
- 45 Van Mierop L H S Pathology and pathogenesis of the common cardiac malformations Cardiovasc Clin 2:27 1970
- 46 Somerville J Atrioventricular defects Mod Concepts Cardiovasc Dis 40:33 1971
- 47 Sakakibara S Yokoyama M Konno H Kudo T and Hatsune K A consideration of operative methods for correction of endocardial cushion defect Ann Thorac Surg 12:419 1971
- 48 Morgan B C Ricketts H J and Wintercheid L C Inferior clockwise frontal plane forces in a child with endocardial cushion defect AM HEART J 82:275 1971
- 49 Baron M G Abnormalities of the mitral valve in endocardial cushion defects Circulation 45:672 1972
- 50 Liberthson R R Paul M H Muster A J Arcilla R A Eckner F A O and Lev M Straddling and displaced atrioventricular orifices and valves with primitive ventricles Circulation 43:213 1971
- 51 Iversen B I Implications of agenesis of the spleen on the pathogenesis of conotruncus anomalies in childhood Acta Paediatr Belg 44 (Suppl 104) 590 1955
- 52 Van Mierop L H S Gesner I H and Schiebler C L Aplasia and polyplasia syndrome Congenital cardiac defects recent advances National Foundation-March of Dimes 8:174 1972

The association of patent ductus arteriosus with CAVO is favorable, and the association with fetal conduction or splenic abnormalities is unfavorable for longevity.

We are indebted to Florence Kotal for her assistance in the preparation of this paper.

REFERENCES

- Gunn F D and Dieckmann J M Malformations of the heart including two cases with common atrioventricular canal and septum defects and one with defect of the atrial septum (cor triatriatum biventriculosum), *Am J Pathol* 3:595 1927
- Rogers H M and Edwards J I Incomplete division of the atrioventricular canal with patent interatrial foramen primum (persistent common atrioventricular ostium) Report of five cases and review of the literature *Am Heart J* 76:28 1948
- Girard G, Latour H, Puech P and Roujon J Les formes anatomiques et les bases du diagnostic de la persistance du canal atriculo-ventriculaire commun *Arch Mal Coeur* 50:909 1957
- Campbell M and Nissen G A H Endocardial cushion defects Common atrioventricular canal and ostium primum *Br Heart J* 19:403 1957
- Haim de Balsac R, Bouchard F, Zilis O, Passelecq J, Guery J, Blonderu P and Dubost C Les Canaux atrio-ventriculaires communs *Bruux Med* 38:325 3:9 and 415 1958
- Kakai C and Edwards J E Pathologic study of persistent common atrioventricular canal *Am Heart J* 56:719 1958
- Paul M H Endocardial cushion defects Persistent common atrioventricular canal and persistent ostium primum *Pediatr Clin North Am* 5:1011 1958
- Dubost C and Blonderu P Canal atrio-ventriculaire et ostium primum Cure chirurgicale d'après 25 observations *J Chir* 78:411 1959
- Girard G, Latour H, Puech P and Oliver G Persistence du canal atriculo-ventriculaire commun II Nouveaux éléments de diagnostic *Montpellier Med* 57:403 1960
- Lev M, Agnasson M H and Arcilla R The pathologic anatomy of common atrioventricular orifice associated with tetralogy of Fallot *Am J Clin Pathol* 36:408 1961
- Molnar M E, Paul M H and Lev M Common A V orifice with pulmonary valvular and hypertrophic subaortic stenosis a case report *Am J Cardiol* 10:291 1962
- Lham J L Persistence du canal Atriculo-ventriculaire commun A propos d'un cas opéré avec succès *Union Med Can* 91:945, 1962
- Van Mierop J H S, Alley R D, Kraus H W and Strimhan A R The anatomy and embryology of endocardial cushion defects *J Thorac Cardiovasc Surg* 43:771 1962
- Manson W C Pathology of the common atrioventricular canal in Morse D P editor *Congenital heart disease* Philadelphia 1967 F A Davis Co p 15 18
- Saint Pierre A, Perrin A, Cahen P and Froment R Persistance du canal atrioventriculaire Commun Complet chez un sujet de seize Ans Réparation Chirurgicale complète mais ictere mortel *Lyon Med* 209:639 1963
- Puech P Les Canaux atrio-ventriculaires embryologiques *Ann Chir Thorac Cardiovasc* 2:734 1963
- Perrin A and Froment R Les formes anatomiques des canaux atrioventriculaires *Ann Chir Thorac Cardiovasc* 11:767 1963
- Young D Criteria for surgery in persistent common atrioventricular canal *Am J Cardiol* 12:80 1963
- Levy M J, Cuello L, Tunst N and Lillehei C W Atrioventricular communis clinical aspects and surgical treatment *Am J Cardiol* 14:587 1964
- Baum D, Roth G J and Creighton S A Right axis deviation, counterclockwise QRS loop and signs of left ventricular underdevelopment in a child with complete type of persistent common atrioventricular canal *Circulation* 30:755 1964
- Kawaji N and Sakamoto T Endocardial cushion defect with transposition of great vessels, pulmonary stenosis and multiple anomalies. A case report *Jpn Heart J* 5:787 1964
- Brown M G, Wolf B S, Steinfield L and Van Mierop L H S Endocardial cushion defects Specific diagnosis by angiocardiology *Am J Cardiol* 13:162 1964
- Gerbode F and Sabar E F Endocardial cushion defects Diagnosis and surgical repair *J Cardiovasc Surg* 5:223 1964
- Sellers R D, Lillehei C W and Edwards J E Subaortic stenosis caused by anomalies of the atrioventricular valves *J Thorac Cardiovasc Surg* 48:289 1964
- Gurod D, Raghib G, Wang Y, Adam P and Amplatz K Angiographic characteristics of persistent common atrioventricular canal *Radiology* 85:442 1965
- Mustard W T, Niguidula F N and Trinkle G A Endocardial cushion defects in infants and children Ten years surgical experience *Br Heart J* 27:768 1965
- Fryer R W M Persistent common atrioventricular canal Anatomy and function in relation to surgical repair *Circulation* 32:170 1965
- Rastelli G C, Kirklin J W and Titus J L Anatomic observations on complete form of persistent common atrioventricular canal with special reference to atrioventricular valve *Mayo Clin Proc* 41:296 1966
- Thomson N B Jr, Niguidula F N and Hohn A Correction of complete atrioventricular canal defect in patients with previous pulmonary artery banding *Am J Cardiol* 18:169 1966
- Lindesmith G G, Meyer H W, Chapman Stanton R E and Jones J C The surgical

Case report

P S, a 5½ year old boy from Italy was well until six months before admission when his mother noted that his heart was beating rapidly. He was asymptomatic except for pallor when examined at that time but his heart rate was found to be 240 and later 180 beats per minute on two separate examination. Treatment with diazepam was started but had no effect. He was hospitalized in Rome and treated sequentially with oral digoxin intravenous digoxin and oral quinidine without response. Next he received an intravenous course of methoxamine and quinidine without improvement. Since his anti-streptolysin titer was noted to be 600 Todd units the diagnosis of rheumatic carditis was considered though there were no clear cut manifestations of rheumatic fever. He received a course of cortisone and digoxin therapy for 20 days. Following this he was treated with a combination of propranolol and digoxin and his heart rate slowed to a constant rate of 150 beats per minute. Three weeks prior to his first evaluation by us he underwent a tonsillectomy for chronic tonsillitis. His heart rate persisted at 150 beats per minute without medication throughout this period.

The patient was first seen by us in April 1972. He presented as a well developed well nourished asymptomatic boy who appeared quite comfortable. Weight 50½ lbs, height 43 inches, pulse 152 and regular, blood pressure 110/80. His chest was symmetrical. The PMI was in the fifth left intercostal space inside the mid clavicular line. There were no murmurs. The ECG showed a coronary sinus rhythm with inverted P waves in Leads 2, 3, and aVF. A 12 hour continuous tape recording of Lead 2 was obtained. This showed persistence of an ectopic atrial focus identified by inverted P waves. The rate was fixed at 152 beats per minute.

He was admitted for further evaluation and possible cardioversion. Serum potassium and thyroxine levels were normal. The hematocrit was 39 per cent, white blood count was 10,000 with a normal differential. At cardiac catheterization his right heart pressures were normal and there was no evidence of an intracardiac shunt. There was a thin radiopaque layer along the right atrial wall that was quite obvious during fluoroscopy. Intra atrial ECG recording showed upright P wave only when the catheter was near the tricuspid valve. Attempts to convert his rhythm to a normal sinus rhythm with intravenous procainamide 200 mg and propranolol 0.5 mg were unsuccessful. Bipolar atrial pacing at a rate of 600 per minute was ineffective. However, after an additional dose of 200 mg of procainamide intravenous bipolar override atrial pacing at 600 per minute for several seconds was followed by conversion to a regular sinus rhythm with a rate of 83 beats per minute for two minutes. A third dose of procainamide (100 mg) resulted in a return to a regular sinus rhythm for 30 minutes. Thereafter, his tachycardia returned and although 500 mg of procainamide, 1 mg of propranolol and 6 mg of dphenhyllantoin were given prior to another attempt at cardioversion, the tachycardia persisted. The patient developed a regular sinus rhythm in association with the vigorous crying that was induced when the pacing catheter

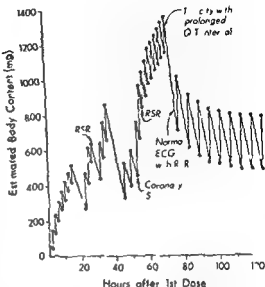


Fig 1 Course of high dosage quinidine therapy. Ordinate represents estimated body content of drug. It was estimated by assuming 100 per cent absorption immediately upon administration and a net half life of eight hours. Regular sinus rhythm (RSR) was present when the level exceeded 450 mg. Toxicity developed when the level exceeded 1300 mg and resolved when it fell to 800 mg. The final regimen (300 mg every six hours) established a plateau of concentration ranges between 500 mg and 800 mg. This succeeded in preventing recurrence of the arrhythmia.

was removed but his atrial tachycardia recurred after two hours of normal rhythm. It continued without change thereafter despite the administration of procainamide 1.5 Gm and propranolol 10 mg for one day, followed by the same dose of propranolol and first 2 Gm and then 3 Gm of procainamide on the two succeeding days. On the third day of this sequence he appeared pale and developed abdominal pain. The ECG continued to show ectopic atrial tachycardia with a rate of 155 beats per minute. The QRS complex, which had measured 0.06 second previously, was prolonged to 0.08 second. Procainamide therapy was discontinued.

We decided to attempt cardioversion with high doses of quinidine (Fig. 1). After a test dose of quinidine a dose of 100 mg was administered every two hours for eight doses without success. The dose was then increased to 200 mg every two hours. After the second dose regular sinus rhythm returned. The next dose was lowered to 100 mg in order to avoid intoxication. However, the coronary sinus rhythm returned two hours later. The following day the 200 mg every two hours regimen was resumed and sinus rhythm returned after the third dose. Since he developed mild hypotension the quinidine was withheld for eight hours. Thereafter the regimen was changed to 200 mg every four hours but this led to return of the arrhythmia. The dose of quinidine was increased to 300 mg every two hours and after the

Drug therapy of heart disease in pediatric patients III. The therapeutic challenge of supraventricular tachyarrhythmias in infants and children

Monika M Rutkowski MD*

Eugene I Doyle, MD**

Sanford N Cohen MD***

New York N Y

Cardiac arrhythmias in the pediatric age group are a serious problem although they occur less frequently than in adults. The incidence of specific types of arrhythmias varies considerably in comparison to adults; for example, atrial flutter and fibrillation are less commonly encountered in children whereas paroxysmal supraventricular tachycardia may occur as frequently in both age groups.

The etiology of arrhythmias in children includes congenital heart disease (e.g. Ebstein's anomaly frequently occurs with supraventricular tachycardias and corrected transposition of the great arteries is frequently associated with atrioventricular block), rheumatic fever, infectious diseases, metabolic diseases, and electrolyte disturbances. Arrhythmias may also be produced by tumors, drugs, mechanical manipulations (e.g. cardiac catheterization) and trauma secondary to open heart surgery. In almost half of the cases, however, no cause can be determined.

Arrhythmias in the pediatric age group may be categorized as follows: (1) the persistent tachyarrhythmias where the atrial and/or ventricular rate is excessive for age; (2) the persistent bradycardias where the ventricular rate is slower than normal and the cardiac output may be inadequate for proper perfusion; and (3) intermittent irregularities (e.g. premature ventricular contractions or premature atrial contractions of less serious physiological consequence).

The only treatment for cardiac arrhythmias prior to several years ago had been the use of pharmacological agents. Recently, the advent of direct current cardioversion and cardiac pacing has broadened our approach to the treatment of the arrhythmias. Most of the arrhythmias seen in infancy and childhood respond to pharmacotherapy, but occasional refractory cases may pose a particularly challenging problem. The following report details an unusually refractory case of tachyarrhythmia in a child.

From the Departments of Pediatrics and Pharmacology, New York University School of Medicine. Received for publication on May 9, 1973.

Reprint requests to: Dr. Monika M. Rutkowski, Department of Pediatrics, New York University School of Medicine, 550 1st Ave., New York, N.Y. 10016.

*Instructor in Pediatrics, New York University School of Medicine.

**Professor of Pediatrics and Director of Pediatric Cardiology, New York University Medical Center.

***Associate Professor of Pediatrics and Pharmacology, New York University School of Medicine. Physician in Charge of Nursery, Bellevue Hospital, John and Mary R. Markle Scholar in Cardiac Medicine.

Table I Antiarrhythmic drugs in pediatric practice

| Drug | Type of arrhythmia | | | | | Contraindications | Dosage† |
|-------------------|--|----------------|---------------------|--------------------------------|--|--|---|
| | Supraventricular tachycardia | Atrial flutter | Atrial fibrillation | Ectopic ventricular arrhythmia | Arrhythmia secondary to digitalis toxicity | | |
| Digoxin | ++ | (+) | ++ | | | Supraventricular tachycardia with AV block ventricular arrhythmia (Use with caution in AV block) | 0.07 mg/Kg < 1 yr 0.06 mg/Kg 1-2 yrs 0.04-0.05 mg/Kg > 2 yrs Total digitalizing dose 1m./po (If i.v. total dose 1/2 of above) |
| Diphenylhydantoin | (+) | | | + | ++ | Use with caution in AV block | 2-3 mg/Kg i.v. |
| Lidocaine | - | - | - | ++ | (+) | Hypotension | 1 mg/Kg i.v. |
| Propranolol | ++ | + | ++ | (+) | (+) | Congestive heart failure AV block obstructive lung disease | 0.05-0.1 mg/Kg i.v. 0.5-1 mg/Kg po (up to 3 m./Kg/day) |
| Procainamide | (+) | (+) | + | ++ | (+) | AV block | 3-5 mg/Kg i.v. |
| Quinidine | + | (+) | ++ | (+) | (+) | AV block | 30 mg/Kg/day po (Usually divided into 5 equal doses) |
| Potassium salt | in cases of low serum potassium concentrations | | | | | | 0.3-0.5 mEq/Kg/hr i.v. |

Symbol ++ = drug of choice + = moderate effect (+) = variable degree of effectiveness - = of little effect
 1) 1 arrhythmic drug m to be given slowly Diphenylhydantoin, lidocaine, propranolol, procainamide, quinidine
 pe ted the same dosage all 20 min ut n.

important to differentiate the therapy of the simple paroxysmal tachycardia in infants and children from the management of the repetitive or persistent tachycardias. In all cases vagal stimulation should be attempted first. This can be achieved through pressure upon either the eyeballs or the area of the carotid sinus. If eyeball pressure is applied care must be taken not to damage the structures themselves. The Valsalva maneuver may be useful in terminating the arrhythmia but is not applicable in most young pediatric patients. The use of other physical maneuvers such as forceful slap on the chest or external cardiac massage may also be successful in some cases.

When physical methods fail digoxin has been the drug of choice for treatment of simple paroxysmal supraventricular tachycardia in infant patients for many years. The majority of patients will eventually respond to digitalis especially if propranolol is added as a second drug. The duration of therapy necessary may vary considerably however. The availability of

direct current countershock as a safe and reliable method¹⁰ to convert the arrhythmia to a normal sinus rhythm without delay has caused us to alter the usual therapeutic recommendation for seriously ill patients.

We withhold digoxin since treatment with direct current countershock may be quite hazardous in the digitalized patient because of the possible excitation of ventricular arrhythmias. We presently attempt cardioversion in these infants first with diphenylhydantoin or propranolol (Tables I and II). Each of these drugs has been used with some success in children with this arrhythmia. The advantage of these agents is that their effect can be seen within minutes and their presence does not contraindicate the use of countershock therapy. If the serum potassium level is low potassium is given (phenylephrine methacholine and neostigmine are rarely used in pediatric patients). If the arrhythmia persists we proceed to direct current countershock. It is our practice to begin with a shock of 10 Watt seconds in young infants

second dose regular sinus rhythm returned. He was maintained with 200 mg of quinidine every two hours thereafter. He developed nausea, vomiting and sinus tachycardia after the eighth dose and quinidine therapy was stopped temporarily.

From this experience it was obvious that it was possible to suppress the ectopic pacemaker with high levels of quinidine and that the therapeutic index (therapeutic dose/toxic dose) was small. Since it was not possible to measure serum quinidine levels, we approximated the biological half life of quinidine in this patient to guide us in calculating an appropriate therapeutic regimen.

We assumed that the half life of quinidine in this child was about eight hours. We estimated that his rhythm became normal when the amount of total body quinidine exceeded 500 mg and that he developed toxic effects when it exceeded 900 mg. We calculated that a dose of 300 mg every 12 hours would keep his total body quinidine between 500 mg and 900 mg. He received quinidine accordingly, to this regimen for the next two days in the hospital and continued to have a normal sinus rhythm without signs of toxicity. He was discharged from the hospital and continued to be treated according to this same quinidine regimen for the next three months with persistence of regular sinus rhythm. The dose was tapered gradually over the following several months. He is presently taking 675 mg per day and is doing well. There are no signs of quinidine toxicity. On rare occasion, especially at times of great excitement (i.e. Christmas day) the patient has experienced short episodes of the tachyarrhythmia. These have responded well to rest and none has lasted more than 30 to 40 minutes.

The etiology for the tachycardia and abnormal pacemaker in this child is obscure. Perhaps a low grade viral pericarditis might have led to both the calcification and the ectopic atrial focus which initiated the arrhythmia. Skin tests for tuberculosis and histoplasmosis were negative. There was no clinical history of an overt pulmonary or pericardial infection in the past.

This case illustrates the difficulties which may be encountered in the treatment of cardiac tachyarrhythmias and also the variety of methods available for their management.

Discussion

There are three types of supraventricular tachycardias which occur in the pediatric age group. The most frequent is the simple paroxysmal supraventricular tachycardia. It occurs most frequently in infancy, especially under four months of age. It is more common in males than in females. It is characterized by a sudden onset and termination, fixed cardiac rates in excess of 180 beats per minute, a brief duration (hours to days) and a prompt response to drug therapy. Among the patients in one large series,¹ the etiology of paroxysmal supraventricular tachycardia during in-

fancy and childhood included Wolff-Parkinson-White syndrome (10 per cent), congenital heart disease (20 per cent), infections, injuries, heart tumors and drugs (20 per cent) and idiopathic (30 per cent). Most cases which occur in infancy are in the idiopathic group. The presenting symptoms in infants most often include pallor, restlessness and poor feeding. This frequently proceeds to frank congestive heart failure, and prompt treatment is mandatory. On the other hand, the older child may present with fever, chest or abdominal pain and palpitations. In most instances the prognosis for infants with supraventricular tachycardia is good and the chance of recurrence decreases considerably after the first year of life. The prognosis for children with supraventricular tachycardia is generally related to the nature of the underlying heart disease.

The second type of supraventricular tachycardia seen among pediatric patients occurs less frequently than the 'simple' type. It is referred to as 'repetitive paroxysmal tachycardia'.^{2,3} This form consists of paroxysms of ectopic beats interrupted by short runs of sinus rhythm. This arrhythmia is resistant to therapy but is not associated with the development of congestive heart failure and therefore carries a favorable prognosis.

The third form of the arrhythmia is encountered very infrequently and is referred to as the sustained ectopic tachycardia or persistent supraventricular tachycardia.^{4,5} It is characterized by the presence of an abnormal P wave which is always easy to discern and by a variable rate. The rate frequently shows diurnal variation. This form of tachycardia usually associated with an ectopic focus may persist for many months or years despite therapy. The prognosis for the majority of children with this type of arrhythmia is good. However, two cases of cerebrovascular accidents and several cases with congestive heart failure have been reported.^{1,4}

The arrhythmia in the patient reported above seemed to be of the third type. In the light of the possibility of complications occurring if the arrhythmia went uncontrolled it was decided to attempt to convert it to regular sinus rhythm despite the fact that he had no symptoms. It is im-

| Inotropic effect | Refractory period duration | | | Toxic effects |
|------------------|----------------------------|------------------------|---------|--|
| | Atrial myocardium | Ventricular myocardium | AV node | |
| — | ↓ | ↓ | ↑ | AV block arrhythmias; tract symptoms |
| 0 | ↓ | ↓ | ↑ | Hypotension |
| 0 | 0 | 0 | 0 | Hypotension convulsions |
| | ↑ | ↑ | ↑ | AV block hypotension |
| (+) | ↑ | ↑ | ↓ | AV block QRS widening hypotension |
| + | ↑ | ↑ | ↓ | QRS widening ventricular arrhythmias hypotension; tract symptoms |

+) = moderate effect 0 = no effect — = negative effect — mod = moderate effect () = minimal negative effect.

way in which it may work is by the interruption of a fixed circus or reciprocating pathway through the atria and especially in the area of the atrioventricular junction.¹² This method has been successful in converting paroxysmal supraventricular tachycardia to a normal sinus rhythm in 70 per cent of adult patients¹² and it is probable that it will have a salutary effect upon the arrhythmia in most infants and children as well.

In treating a child with the sustained or ectopic form of supraventricular tachycardia it is possible that one will have to be satisfied with the control of the ventricular rate without conversion of the atrial arrhythmia. It is generally considered a matter of personal preference as to which sequence of pharmacological agents is followed. We prefer to start with diphenylhydantoin because it has a minimal negative inotropic effect and acts within minutes. We prefer to use the beta-adrenergic blocking agent propranolol as the second drug in the treatment of such patients. Its antiarrhythmic mechanism is related to

both its adrenergic blocking effect and its ability to depress conductivity in a quinidine-like manner. This drug should be used with caution in cases with severe congestive heart failure and is contraindicated in patients with atrioventricular block or obstructive lung disease.

Procainamide may be useful in the treatment of refractory cases of supraventricular tachycardia but it is generally more effective in the treatment of ventricular arrhythmias. Digoxin (or other digitalis glycosides) is the final drug on our usual list. It acts by increasing the refractory period of the atrioventricular node and thereby slows the ventricular rate. In some cases it acts to abolish the arrhythmia itself. More often quinidine is required as an additional drug to suppress the ectopic pacemaker in these complicated cases.

All of these drugs were used in our patient at one time or another during his hospital course. However, they did not alter the activity of the ectopic pacemaker. Following the use of pharmacotherapy it was decided to attempt atrial pacing in this

Table II Mechanism of action and toxic effects of antiarrhythmic drugs*

| Drug | Purkinje fibers | | Automaticity of | | | Intoxi- cated |
|----------------------|----------------------------|---|--------------------|--------------------------------|-------------------------------------|------------------|
| | action potential duration | | Purkinje fibers | Atrial myocardial fibers | Ventricular myocardial fibers | |
| | refractory period duration | | | | | |
| Digitalis glycosides | | | ↑ | ↑ | ↑ | ++ |
| Diphenylhydantoin | ↓ | ↓ | ↓ | ↓ | ↓ | (-) |
| Lidocaine | ↓ | ↓ | ↓ | unknown | ↓ | () |
| Propranolol | ↓ | ↓ | ↓ | ↓ | ↓ | -- |
| Procainamide | ↑ | ↑ | ↓ | ↓ | ↓ | - |
| Quinidine | ↑ | ↑ | ↓ | ↓ | ↓ | -- |

*Symbols ↑ = increase ↓ = decrease } amplitude of arrow indicates relative degree of effect ++ = strong positive effect + = moderate positive effect

and to slowly increase this until conversion occurs. It is rare to require more than 25 Watt seconds in these patients. It is not necessary to administer general anesthesia or muscle relaxants to these patients since there is no generalized convulsive effect of this small shock. In some cases meperidine 1 mg per kilogram of body weight or diazepam 2 to 3 mg may be given for analgesia or relaxation. Once there has been a return to normal cardiac rhythm we digitalize our patients and maintain them on digoxin for 3 to 6 months since recurrences of the arrhythmia are frequent when it begins in the first year of life.

Children with the simple form of paroxysmal supraventricular tachycardia rarely present with critical illness accompanied by congestive heart failure. They are generally treated in sequence with diphenylhydantoin, propranolol, procainamide and digoxin. In some cases only a combination of several of these drugs (e.g. digoxin and propranolol) will be successful. If none of these agents or combinations is effective, we prefer to use transvenous atrial

pacing rather than direct current counter shock as the next step in therapy. Some investigators feel that both of these procedures carry less risk than quinidine therapy.¹¹ The advantages of atrial pacing over the direct current countershock method are that there is no need for discontinuation of digoxin therapy and there is no need for general anesthesia. A transvenous bipolar catheter electrode can be inserted percutaneously without trouble and has the advantage that it can be left in place in case of frequent recurrences of the arrhythmia. The mechanism of atrial pacing and the effect of such a procedure upon the tachycardia have been reviewed in detail elsewhere.¹²⁻¹⁴

Atrial pacing may convert a supraventricular tachycardia to a normal sinus rhythm by three different mechanisms. It may suppress an ectopic pacemaker through overdrive. It may convert a self-sustained tachycardia such as atrial flutter into one which is not sustained such as fibrillation which results in a return to sinus rhythm in many cases. The final

| Effect | Refractory period duration | | | Toxic effects |
|--------|----------------------------|------------------------|---------|--|
| | Atrial myocardium | Ventricular myocardium | AV node | |
| | ↓ | ↓ | ↑ | AV block arrhythmias g: tract symptoms |
| 0 | ↓ | ↓ | ↑ | Hypotension |
| 0 | 0 | 0 | 0 | Hypotension convulsions |
| - | ↑ | ↑ | ↑ | AV block hypotension |
| (+) | ↑ | ↑ | ↓ | AV block QRS widening hypotension |
| + | ↑ | ↑ | ↓ | QRS widening ventricular arrhythmias hypotension g: tract symptoms |

0 = no effect 1 = slight effect 2 = moderate effect 3 = marked effect 4 = severe effect 5 = fatal effect

way in which it may work is by the interruption of a fixed circus or reciprocating pathway through the atria and especially in the area of the atrioventricular junction.¹² This method has been successful in converting paroxysmal supraventricular tachycardia to a normal sinus rhythm in 70 per cent of adult patients¹³ and it is probable that it will have a salutary effect upon the arrhythmia in most infants and children as well.

In treating a child with the sustained or ectopic form of supraventricular tachycardia it is possible that one will have to be satisfied with the control of the ventricular rate without conversion of the atrial arrhythmia. It is generally considered a matter of personal preference as to which sequence of pharmacological agents is followed. We prefer to start with diphenylhydantoin because it has a minimal negative inotropic effect and acts within minutes. We prefer to use the beta-adrenergic blocking agent propranolol as the second drug in the treatment of such patients. Its antiarrhythmic mechanism is related to

both its adrenergic blocking effect and its ability to depress conductivity in a quinidine-like manner. This drug should be used with caution in cases with severe congestive heart failure and is contraindicated in patients with atrioventricular block or obstructive lung disease.

Procainamide may be useful in the treatment of refractory cases of supraventricular tachycardia but it is generally more effective in the treatment of ventricular arrhythmias. Digoxin (or other digitalis glycosides) is the final drug on our usual list. It acts by increasing the refractory period of the atrioventricular node and thereby slows the ventricular rate. In some cases it acts to abolish the arrhythmia itself. More often quinidine is required as an additional drug to suppress the ectopic pacemaker in these complicated cases.

All of these drugs were used in our patient at one time or another during his hospital course. However, they did not alter the activity of the ectopic pacemaker. Following the use of pharmacotherapy it was decided to attempt atrial pacing in this

patient. The atrial "overdrive" succeeded in returning him to a regular sinus rhythm only when it was combined with procainamide for the suppression of the ectopic focus. The sinus rhythm was not stable, however, and he reverted to his arrhythmia soon thereafter.

We decided to make a final attempt to convert his arrhythmia to a normal sinus rhythm, this time with high doses of quinidine. High dosage quinidine therapy can be extremely hazardous and should be carried out only in an intensive care unit that is equipped and staffed for patient monitoring and resuscitation. The use of serum quinidine concentration measurements to guide therapy does not eliminate the hazard of such a regimen since therapeutic effect and/or toxicity may not be predictable on the basis of serum level alone. In our patient we assumed that the drug was absorbed completely as soon as a dose was given and that its half life in the body was 8 hours. This approach permitted us to estimate the proper therapeutic regimen for him once the therapeutic range had been defined (Fig 1).

The authors are indebted to Miss Lucille M. Rung and Mrs. Leanne Rappaport for the preparation of these manuscripts.

REFERENCES

- Nadas A S and Fyler D C. *Pediatric cardiology*. 3rd ed. Philadelphia: London and Toronto 1972. W B Saunders Company.
- Parkinson J and Papp C. Repetitive paroxysmal tachycardia. *Br Heart J* 9:241 1947.
- Morgan C L and Nadas A S. Chronic ectopic tachycardia in infancy and childhood. *Am Heart J* 68:617 1964.
- Shrout N, Spellman S and Rubin I. Persistent supraventricular tachycardia. Case report with review of literature. *Circulation* 10:232 1954.
- Hay J D and Keidan S E. Persistent ectopic auricular tachycardia in children. *Br Heart J* 14:345 1952.
- Lown B. Electrical reversion of cardiac arrhythmias. *Br Heart J* 29:469 1967.
- Paul M H and Miller R A. External electrical termination of supraventricular arrhythmias in congenital heart disease. *Circulation* 25:604 1962.
- Pryor R and Blount G S Jr. Refractory supraventricular tachycardia in infancy. *Am J Dis Child* 107:428 1964.
- White R I and Humphries J. Direct current electroshock in the treatment of supraventricular arrhythmias. *J Pediatr* 70:119 1967.
- Hessneruck A, Chojnacki H and Barker H J. Cardioversion of auricular flutter in a newborn infant. *Am J Cardiol* 15:726 1965.
- Chou T. Treatment of cardiac arrhythmias. *Mod Treat* 7:40 1970.
- Barold S S and Linhart J W. Recent advances in the treatment of ectopic tachycardias by electrical pacing. *Am J Cardiol* 23:698 1970.
- Cheng T O. Atrial pacing: its diagnostic and therapeutic applications (Review). *Progr Cardiovasc Dis* 14:230 1971.
- Hunsaker M R and Khoury G H. Management of supraventricular tachycardia by atrial stimulation. *J Pediatr* 77:454 1970.

1 Nadas A S and Fyler D C. *Pediatric cardiology*. 3rd ed. Philadelphia: London and

Theoretic considerations of the post-systolic "dip" of constrictive pericarditis

Following its original description in 1946¹ various explanations have been offered for the presence of the post systolic dip (Fig 1) seen in pressure tracings recorded from the cavity of the right ventricle of patients with chronic constrictive pericarditis. It has been suggested that the dip is an artifact produced by a low frequency recording apparatus² but the pattern has been found consistently in patients with constrictive pericarditis since high frequency recording equipment has become available. Another explanation for the presence of the diastolic dip is that it represents an actual cardiodynamic event^{3,4,5} because the ventricles are completely empty at the end of systole and the atrial pressure is high. Rapid ventricular filling takes place which is abruptly terminated by the thickened pericardium. However this theory does not explain why the ventricular relaxation is not isometric with time.

Although the possibility of elastic recoil of the thickened pericardium and chest wall has been mentioned by some⁶ a precise mechanism has not been offered. We believe that a sudden rather than abrupt elastic recoil of the distorted (during systole) thickened pericardium when the ventricle rather suddenly relaxes is responsible for the diastolic dip in constrictive pericarditis and we therefore offer the following hypothesis.

The heart in constrictive pericarditis is encased in a thick shell of fibrous connective tissue with or without calcium which is fixed to and therefore a part of the ventricular wall. Thus when the ventricle contracts work is performed on this thick wall which deforms in much the same manner as when one loads a spring. Following termination of myocardial contraction and with the sudden unloading of the distorted fibrous pericardium with the onset of diastole the ventricular wall abruptly springs back into place suddenly reducing intracavitary pressure thereby literally sucking blood from the atrium into the ventricle. This recoil of the ventricular wall results in the rapid fall in ventricular pressure which is responsible for the ventricular dip as recorded in the intracavitary pressure tracing. The ventricle fills rapidly and therefore an early plateau of the right ventricular pressure tracing is seen (Fig 1).

Supporting evidence for this concept is seen in experimental cardiac tamponade produced with air or fluid. In such a situation the fluid or air in the pericardial cavity is not fixed to the myocardial wall even though diastolic filling of the ventricle is re-

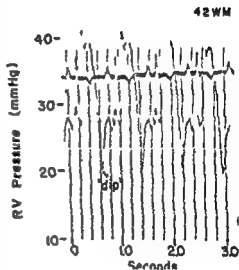


Fig 1 Time course of right ventricular pressure showing post-systolic dip in constrictive pericarditis

stricted. Thus during systole there is no physiologic spin to load nor is there one to suddenly unload during the early phase of diastole. Under such circumstances therefore the characteristic dip in right ventricular pressure is not expected nor recorded.⁷

Parenthetically it may be noted that the work performed in deforming the pericardium in constrictive pericarditis is not useful work and therefore myocardial contraction is less efficient and work and energy are wasted loading a physiologic spring. This wasted work must contribute in part to the extra work of the heart which in turn contributes to early myocardial failure observed in patients with constrictive pericarditis.

A possible comparison can be made between the heart in constrictive pericarditis and a heart with myocardial fibrosis in which the myocardium has been partially replaced by fibrous connective tissue. In the latter heart the amount of contractile element has been reduced and/or the remainder diseased and in addition unnecessary work by an injured myocardium is required to deform the interstitial fibrous connective tissue. The spring load concept is reflected in the right ventricular pressure

tracings from patients with such hearts which are similar to the tracings recorded from patients with constrictive pericarditis.⁸ The same concepts apply to hearts with endocardial fibrosis and thickening.

C E Burch MD

T D Giles MD

Department of Medicine

Tulane University School of Medicine

1430 Tulane Ave

New Orleans La 70112

REFERENCES

- Bloomfield R A, Larson H D, Courmand A, Breed F S and Richards D W Jr. Recording of right heart pressures in normal subjects and in patients with chronic pulmonary disease and various types of cardio-circulatory disease. *J Clin Invest* 25:639 1946
- Wiggers C J. Physiology in health and disease. Philadelphia 1949 Lea & Febiger p 648
- McKusick V A. Chronic constrictive pericarditis. *Bull Johns Hopkins Hosp* 90:3 1952
- Hansen A T, Eskildsen P and Gotzsche H. Pressure curves from the right atricle and the right ventricle in chronic constrictive pericarditis. *Circulation* 3:881 1951
- Harvey R M, Ferrer M I, Cathcart R T, Richards D W and Courmand A. Mechanical and myocardial factors in chronic constrictive pericarditis. *Circulation* 31:695 1953
- Yu P N G, Lovejoy I W Jr, Joos H A, Nye R E Jr and Mahoney E B. Right atricular and ventricular pressure patterns in constrictive pericarditis. *Circulation* 7:102 1953
- Fowler N O, Shabetai R and Braunstein J R. Transmural ventricular pressures in experimental cardiac tamponade. *Circ Res* 7:733 1959
- Somers K, Brenton D P, Darbela P G, Fowler J M, Kanyerezi H R and Sood N H. Hemodynamic features of severe endomyocardial fibrosis of right ventricle including comparison with constrictive pericarditis. *Br Heart J* 30:322 1968

Blood flow in the internal mammary artery

Use of the internal mammary artery (IMA) as a coronary bypass graft has been criticized on the basis of its small caliber and limited flow capacity. However, a late failure rate as high as 30 per cent has been reported for saphenous vein in the first year^{1,2} in contrast to three per cent for the IMA.³ Therefore, in the past 10 months the IMA has been used 78 times for coronary bypass grafting.

In most instances free graft flow was measured by a volumetric timed collection immediately prior to making the anastomosis. After discontinuation of cardiopulmonary bypass, IMA flow was measured using an electromagnetic flow probe. Observations were made of basal flow, of reactive hyperemia flow

after a 30 second graft occlusion and of peak flow after injection of 15 mg of papaverine into the graft. (Papaverine injection was performed in the first 34 grafts and then discontinued.) This data is compared with flow data obtained in the standard manner from a much larger series of saphenous vein coronary bypass grafts. Free flow was not measured in the vein grafts (see Table I).

It is apparent that basal graft flows are similar for the IMA and saphenous vein whether to the right coronary artery (RCA) or to the left anterior descending artery (LAD). Both reactive hyperemia flow and peak flow are significantly less for the IMA. These observations indicate that the flow

Table I. Flow data for internal mammary artery bypass grafts compared to saphenous vein coronary bypass grafts

| Variables | Right IMA* | Left IMA | Vein RCA | Vein LAD |
|--------------------------------|------------|----------|----------|----------|
| Number of grafts | 17 | 51 | 185 | 213 |
| Free flow (ml/min) | 86 ± 9 | 98 ± 6 | — | — |
| Mean arterial pressure (mm Hg) | 81 ± 4 | 93 ± 3 | — | — |
| Basal flow | 68 ± 9 | 56 ± 4 | 61 ± 4 | 57 ± 3 |
| Reactive hyperemia | 68 ± 9 | 59 ± 4 | 86 ± 4 | 75 ± 4 |
| Peak flow (papaverine) | 79 ± 7 | 97 ± 9 | 174 ± 8 | 155 ± 7 |

*Abbreviations: IMA = internal mammary artery graft; RCA = right coronary artery graft; LAD = left anterior descending artery graft.

capacity of the IMA is limited and may not meet myocardial requirements 1 times of maximal need. However the IMA is probably not at its ultimate caliber during the flow measurements. Without question the IMA develops spasm during mobilization and antispasmodic agents such as Nylomax or papaverine have not been used to remedy this. Initially gentle balloon catheter dilation was used but this resulted in intimal fracture in one graft and this maneuver was discontinued. In three IMA grafts flow was remeasured 1x to eight hours later at the time of reoperation for postoperative hemorrhage. At reoperation basal flow was 40 per cent greater than previously. Vein graft flow has not been greater at reoperation. This would indicate that the IMA dilates in the hours following operation and its flow capacity increases. The luminal diameter of the IMA has been 1.3 or 2.0 mm as measured with malleable probes in 0.5 mm gradations. In only one instance was the IMA not used because of inadequate free flow and in this instance a hemostatic clip had been inadvertently placed across the artery at the first intercostal space.

In the final analysis the apparent better long

term patency of the IMA must be weighed against its limited flow capacity and the technical difficulties attending its use. It is important to realize that once the sternum has been closed the IMA is lost since sternal closure may directly destroy it or secondary cicatrix precludes its future mobilization. Therefore the IMA cannot be held in reserve should saphenous vein coronary grafts fail but the IMA must be utilized at the initial operation if it is to be used at all.

Hendrick B. Barner, MD
Department of Surgery
St. Louis University
St. Louis, Mo 63104

REFERENCES

1. Bourassa M, C. Lesperance J, Campeau L, and Simard P. Factors influencing patency of aortocoronary vein grafts. *Circulation* 45 (Suppl 1): 19 1972.
2. Green G E. Internal mammary artery coronary artery anastomosis: three year experience with 165 patients. *Ann Thorac Surg* 14: 260 1972.

Cell growth in man

If the object of the process of education is to equip a child for adult life, so the practice of pediatrics and especially the conduct of research in pediatrics must be chiefly concerned with the long term effects of disease processes. The prime function of the pediatrician is to produce a healthy adult. On the other hand diseases which usually present in adult life probably have little chance of improvement if treatment is not instituted early in life. The long term treatment of males with Type 2 hyperlipoproteinemia may be taken as an example of such a condition. In childhood the basis of many pathogenetic processes is laid and environmental circumstances which impinge on the early life of a child may well carry life-long consequences.

The effect of the environment on a developing organism depends largely on timing. Thus the effects of rubella on the eight week-old fetus are catastrophic, while they are negligible on an eight year old child. It is probable that the same principle applies to all infants. It is possible for example to alter growth processes in animals experimentally simply by over- or under-nutrition¹ and in general the earlier the treatment the more severe its effect. Much information on nutrition on the brain growth of animals on these lines has accumulated and the indication is that undernutrition at certain critical periods can permanently reduce the number of brain cells if the treatment is of sufficient severity and duration. Similar experiments have shown that mental development may also be impaired.

Experiments such as these are clearly not possible

in man. It is known that head size can be affected by undernutrition during the period of most rapid brain growth in utero in the perinatal period² and in early childhood.³ In the latter study the reduction in head size of children dying of marasmus before the age of one year was directly related to the decrease in the number of brain cells. Because of the difficulty in obtaining such data, information hitherto available in man is scanty to say the least and it is not always justifiable to extrapolate directly from animal experiments to the human situation.

Body fat is unique as an organ in man in being easily sampled and relatively easily measured from skinfold thicknesses. It appears in the fetus late in gestation (at about 30 weeks) and its period of most rapid growth continues during the first year of postnatal life. In this organ it is therefore possible to study directly the effects of different environmental circumstances occurring in a period which is usually relatively well documented.

I have studied the growth in the number of adipose cells in children to whom a series of environmental accidents have occurred. Control values were obtained from otherwise healthy children undergoing elective surgery; the other groups comprised one of obese children, half of whom had gained weight excessively in the first year of life and half had not; one of children with growth hormone deficiency either idiopathic or following surgery for craniopharyngiomas and one of children whose birthweight was at least 2 SD below the mean after the length of gestation had been allowed for.

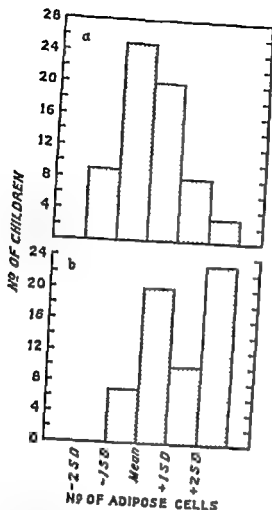


Fig 1 Total number of adipose cells Panel a control children Panel b obese children

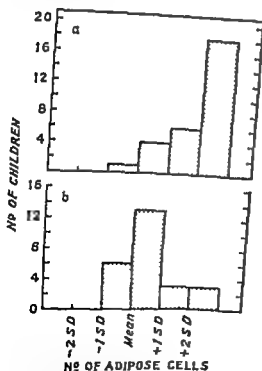


Fig 2 Effect of age at onset of obesity on the number of adipose cells in obese children Panel a early onset obese children Panel b late onset obese children

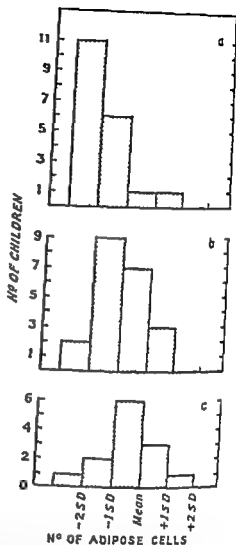


Fig 3 Effect of birth weight and growth hormone deficiency on the number of adipose cells in children Panel a light for their age children Panel b children with idiopathic isolated growth hormone deficiency Panel c children growth hormone deficient due to surgery for craniopharyngiomas

The number of adipose cells in obese children was increased (Fig 1) but the increase was entirely accounted for by the contribution made by the early obese children. The later obese children though they were of similar fatness had a number of adipose cells which did not differ from control (Fig 2). Similar findings applied to a group of obese adults whom I have studied.⁷ Whether to have an increased number of adipose cells matters is not known but in view of the mortality rate and the morbidity associated with the condition it is a question which clearly demands an answer. Since rapidly gaining babies also have an increased incidence of obesity later,⁸ it is more than of passing importance because it may be that we should reconsider our whole practice of infant feeding.

In the growth hormone deficient and light for their age children the situation was the reverse (Fig 3). Light for their age children suffer a severe insult to adipose cell division early in the period of adipose organogenesis and have a profound deficit in cell number. In children with idiopathic isolated

growth hormone deficiency the lack becomes operative presumably at birth later in organogenesis and the cell deficit is less. Children who lack growth hormone at a later stage would be expected to have passed the age when the adipose organ is sensitive to such circumstances. These children would be expected to have a normal complement of cells when the growth delay induced by growth hormone deficiency and evidenced by skeletal retardation is allowed for. The findings in the group of children with craniopharyngiomas confirm this expectation.

Thus the adipose organ it seems has a finite and well-defined sensitive period during which external circumstances have lasting effects on cellular growth. Such a model presumably applies with different timings to all organs in man.

C G D Brook MD MRCP
Lecturer in Child Health and Growth
Institute of Child Health
London WC1N 1EH England
Present address
Department of Endocrinology
Kandervall
CH 8032 Zurich Switzerland

REFERENCES

Widdowson E M and McCance R A. Some effects of accelerating growth. *Proc. R. Soc. Lond. (Biol.)* 152:188 1960.

- 2 Widdowson E M and McCance R A. The effect of finite periods of undernutrition at different ages on the composition and subsequent development of the rat. *Proc. R. Soc. Lond. (Biol.)* 158:329 1963.
- 3 Zamenhof S, Van Marthens E and Margolis F L. DNA and protein in neonatal brain after atrophy by maternal dietary protein restriction. *Science* 160:3825 1968.
- 4 Simonson M and Cjow B F. Maze studies on progeny of underfed mother rats. *J. Nutr.* 100:685 1970.
- 5 Davies P A and Davis J P. Very low birth weight and subsequent head growth. *Lancet* 2:1216 1970.
- 6 Winick M and Rosso P. The effect of severe early malnutrition on cellular growth of the human brain. *Pediatr. Res.* 3:181 1969.
- 7 Brook C G D, Lloyd J K and Wolff O H. Relation between age of onset of obesity and size and number of adipose cells. *Br. Med. J.* 2:25 1972.
- 8 Eid E E. Follow up study of physical growth of children who had excessive weight gain in the first six months of life. *Br. Med. J.* 2:174 1970.

Serum enzyme levels after operation

The differential diagnosis of chest pain after operation is complicated by the fact that those serum enzymes which are characteristically elevated after myocardial infarction¹ (and to a lesser extent after pulmonary infarction²) are also often raised postoperatively.³ Therefore in order to establish the duration and extent of the serum enzyme level changes postoperatively serial measurements of a pyruvate transaminase (AsT), hydroxybutyric dehydrogenase (HBD) and creatine kinase (CK) levels were made after a variety of cold operations.⁴

Fourteen patients undergoing thoracic or abdominal operations were studied, having excluded those in whom the enzymes were elevated preoperatively, those with abnormal electrocardiograms and those in whom there was any evidence of deep vein thrombosis or myocardial or pulmonary infarction. Of the 14 patients, four underwent thoracotomy, three cholecystectomy, three vagotomy and pyloroplasty, two partial gastrectomy, one colectomy and one suprapubic prostatectomy. The AsT and HBD levels were measured by spectrophotometric methods (normal ranges 4 to 15 and 50 to 150 IU per liter respectively) and the CK levels by a colorimetric method (normal range 4 to 60 IU per liter). The enzyme levels were

measured preoperatively and at 24 hours, three, five, seven and 10 days postoperatively.

In 10 patients there was an elevation of all three enzymes, maximum levels being attained at 24 hours or three days after which they gradually fell to normal levels. The highest levels recorded were AsT 62 IU per liter, HBD 406 IU per liter and CK 550 IU per liter. These figures are within the range of enzyme results frequently encountered after myocardial infarction.¹ In every case in this series the HBD had returned to normal within five days of operation, the CK within seven days and the AsT was normal in most patients by 10 days at which time all AsT levels were below 20 IU per liter.

The rises in serum enzyme levels after operation are probably due to trauma to tissues, especially skeletal muscle.⁵ This is supported by the finding that the highest enzyme rises occurred after operations involving the most extensive operative trauma (thoracotomy patients).³

Elevated serum enzyme levels after operation must therefore be interpreted with extreme caution. After the fifth postoperative day the significance of raised serum enzyme levels increases with respect to the diagnosis of postoperative myocardial or pulmonary infarction. Elevated levels after the tenth

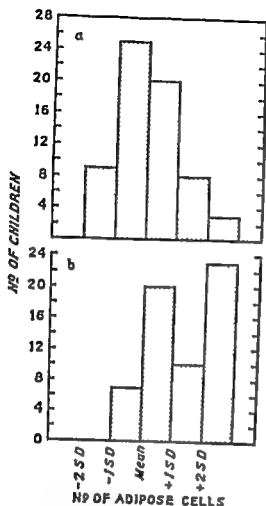


Fig 1 Total number of adipose cells Panel a control children Panel b obese children

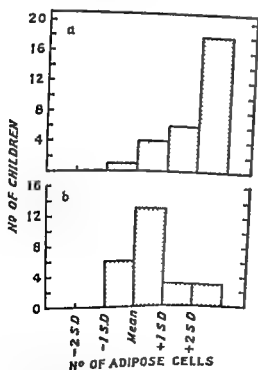


Fig 2 Effect of age at onset of obesity on the number of adipose cells in obese children Panel a early onset children Panel b late onset children

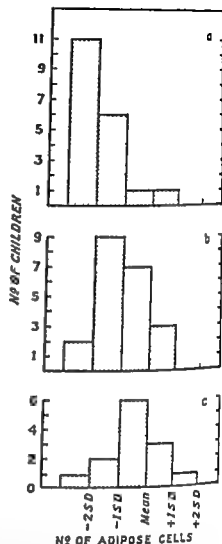


Fig 3 Effect of birth weight and growth hormone deficiency on the number of adipose cells in children Panel a light for their age children Panel b children with idiopathic isolated growth hormone deficiency Panel c children growth hormone-deficient due to surgery for craniopharyngiomas.

The number of adipose cells in obese children was increased (Fig 1) but the increase was entirely accounted for by the contribution made by the early onset obese children. The later onset children (those who were of similar fatness) had a number of adipose cells which did not differ from control (Fig 2). Similar findings applied to a group of obese adults whom I have studied.⁷ Whether to have an increased number of adipose cells matters is not known, but in view of the mortality rate and the morbidity associated with the condition it is a question which clearly demands an answer. Since rapidly gaining babies also have an increased incidence of obesity later,⁸ it is more than of passing importance because it may be that we should reconsider our whole practice of infant feeding.

In the growth hormone deficient and light for their age children the situation was the reverse (Fig 3). Light for their age children suffer a severe insult to adipose cell division early in the period of adipose organogenesis and have a profound deficit in cell number. In children with idiopathic isolated

growth hormone deficiency the lack becomes operative presumably at birth, later in organogenesis and the cell deficit is less. Children who lack growth hormone at a later stage would be expected to have passed the age when the adipose organ is sensitive to such circumstances. These children would be expected to have a normal complement of cells when the growth delay induced by growth hormone deficiency and evidenced by skeletal retardation is allowed for. The findings in the group of children with craniopharyngiomas confirm this expectation.

Thus the adipose organ it seems has a finite and well-defined sensitive period during which external circumstances have lasting effects on cellular growth such a model presumably applies with different timings to all organs in man.

C G D Brook MD FRCP
Lecturer in Child Health and Growth
Institute of Child Health
London WC1N 1EH England
Present address
Department of Endocrinology
Aandspital
CH 8032 Zurich Switzerland

REFERENCES

- 1 Widdowson E M and McCance R A Some effects of accelerating growth Proc R Soc. Lond (Biol.) 152:188 1960

- 2 Widdowson E M and McCance R A The effect of finite periods of undernutrition at different ages on the composition and subsequent development of the rat Proc. R Soc. Lond (Biol.) 153:329 1963
- 3 Zamenhof E Van Marthens E. and Margolis F L. DNA and protein in neonatal brain after ation by maternal dietary protein restriction Science 160:3825 1968
- 4 Simpson M and Cjow B F Maze studies on progeny of underfed mother rats J Nutr 100 685 1970
- 5 Davies P A and Davis J P Very low birth weight and subsequent head growth Lancet 2 1216 1970
- 6 Winick M and Rosso P The effect of severe early malnutrition on cellular growth of the human brain Pediatr Res 3:181 1969
- 7 Brook C G D Lloyd J K. and Wolff O H Relation between age of onset of obesity and size and number of adipose cells Br Med J 2:25 1972
- 8 Eid E E Follow up study of physical growth of children who had excessive weight gain in the first six months of life Br Med J 2:74 1970

Serum enzyme levels after operation

The differential diagnosis of chest pain after operation is complicated by the fact that those serum enzymes which are characteristically elevated after myocardial infarction¹ (and to a lesser extent after pulmonary infarction) are also often raised postoperatively.² Therefore in order to establish the duration and extent of the serum enzyme level changes postoperatively serial measurements of a pyruvate transaminase (AsT) hydroxybutyrate dehydrogenase (HBD) and creatine kinase (CK) levels were made after a variety of cold operations.³

Fourteen patients undergoing thoracic or abdominal operations were studied having excluded those in whom the enzymes were elevated preoperatively those with abnormal electrocardiograms and those in whom there was any evidence of deep vein thrombosis or myocardial or pulmonary infarction. Of the 14 patients four underwent thoracotomy three cholecystectomy three vagotomy and pyloroplasty two partial gastrectomy one colectomy and one suprapubic prostatectomy. The AsT and HBD levels were measured by spectrophotometric methods (normal ranges 4 to 15 and 50 to 150 I U per liter respectively) and the CK levels by a colorimetric method (normal range 4 to 60 I U per liter). The enzyme levels were

measured preoperatively and at 24 hours three five seven and 10 days postoperatively.

In 10 patients there was an elevation of all three enzymes maximum levels being attained at 24 hours or three days after which they gradually fell to normal levels. The highest levels recorded were AsT 62 I U per liter HBD 406 I U per liter and CK 550 I U per liter. These figures are within the range of enzyme results frequently encountered after myocardial infarction.¹ In every case in this series the HBD had returned to normal within five days of operation the CK within seven days and the AsT was normal in most patients by 10 days at which time all AsT levels were below 20 I U per liter.

The rises in serum enzyme levels after operation are probably due to trauma to tissues especially skeletal muscle.⁴ This is supported by the finding that the highest enzyme rises occurred after operations involving the most extensive operative trauma (thoracotomy patients).⁵

Elevated serum enzyme levels after operation must therefore be interpreted with extreme caution. After the fifth postoperative day the significance of raised serum enzyme levels increases with respect to the diagnosis of postoperative myocardial or pulmonary infarction. Elevated levels after the tenth

postoperative day can no longer be attributed to operative trauma

Sylvia M Watkins B M M R C P
Adam Lewis F R C S
The Royal Free Hospital
North Western Branch
Lawn Rd
London N W 3 England

REFERENCES

- 1 Henley K E Schmidt E and Schmidt F W
Enzymes in serum Springfield Ill 1966
Charles C Thomas Publisher
- 2 Watkins S M and Lewis A Serum enzyme
levels in the diagnosis of postoperative myo-
cardial infarction *Br Med J* 3:733 1972

Book reviews

SCLERODERMA Proceeding of an International Symposium sponsored by the World Health Organization Edited by Prof F Delbarre Paris 1972 Masson et Cie 311 pp

This series of papers represent the proceedings of a symposium sponsored by the World Health Organization. The papers are in French or English. This book is concerned with seven main subjects namely general aspects of scleroderma scleroderma as a cutaneous disease scleroderma as a systematic disease the clinical forms diagnostic problem anatomic and pathologic changes and treatment. In essence this is a summary of the practical clinical problems of scleroderma a somewhat rare but important disease. This review is a good one and the book certainly is useful in that it contains the important aspects of the problems of scleroderma presented for the clinician. Doctors will find this to be an excellent review of the subject.

ADVANCES IN CARDIOLOGY Early Diagnosis of Coronary Heart Disease P Halonen and A Loubja Editors Helsinki Basel Munchen Paris London New York Sydney 1973 S Karger AG 231 pp Price \$24.20

This important book describes the present day idea on the early diagnosis of coronary heart disease. It really contains nothing especially new but it brings together for readers the concepts of clinicians concerning their own experience with the early diagnosis of coronary heart disease. The book is certainly worth owning and studying. Coronary arteriography emotional factors hyperlipemia apexcardiography atrial pacing measurements of coronary flow prevention and early diagnosis are among the many subjects discussed. This is a useful book for valuable reading by those with limited time.

BLOOD FLOW MEASUREMENT Edited by Colin Roberts Baltimore 1972 The Williams & Wilkins Company 184 pp Price \$18.75

Dr Roberts has edited a book which is concerned briefly with aspects of the methods and techniques for measuring blood flow in the heart and large blood vessels. The peripheral flow such as in the digits and microcirculation is neglected. Each presentation of which there are 34 in 181 pages can attest to the brevity of each discussion. Nevertheless those who are seriously engaged in studying and measuring blood flow will find this book interesting. The problems difficulties and inaccuracies of the various techniques are not criti-

cally discussed. This would not be expected in this small book. The book is well written and clearly presented so that readers can learn the methods and principles involved in measuring blood flow in large vessels.

ANTIARRHYTHMIC AGENTS Arthur J Moss M.D. and Robert H Patton M.D. Springfield Ill 1973 Charles C Thomas Publisher 161 pp

Moss and Patton have produced a book which is concerned primarily with ten antiarrhythmic agents commonly used in medicine at present. The electrophysiology of the heart is briefly described and the pharmacology and clinical use of the antiarrhythmic agents presented in separate chapters. The entire book contains only 112 pages and therefore would not be expected to contain detailed information about these drugs. The authors have presented their own concepts in management briefly. Readers will find this to be a short quick source of data on important drugs.

MYOCARDIAL BLOOD FLOW IN MAN Methods and Significance in Coronary Disease A Maseri M.D. Editor Turin 1972 Minerva Medica 576 pp Price \$35.00

This book contains the papers and discussions on Myocardial Blood Flow in Man presented at a meeting in Pisa during June 10 to 12 1971. The participants were from many nations and the subjects discussed were quite varied. The book is divided into six parts and a round table discussion. The six parts were concerned with determinants of myocardial blood flow transmural distribution theoretical and technical problems hemodynamic correlates significance of measurements adaptations to ischemia and adaptations of myocardial perfusion to coronary artery disease. Discussions follow most of the papers. A considerable number of problems are discussed. They are of importance and should interest physiologists and cardiologists since ischemic heart disease is so common. In spite of the extensive studies in progress it is still not possible to measure quantitatively the rate of coronary blood flow with sufficient accuracy and sensitivity to meet the needs required to supplement the clinical data already obtainable. The procedures available for intact man are too complex and insensitive to be of practical value. This is evident from this book. Nevertheless this is an important publication which should interest cardiologists and physiologists. The organizing group for the symposium has done a fine service to cardiology.

Announcements

American Institute of Ultrasound Meeting

The Eighteenth Annual Meeting, of The American Institute of Ultrasound in Medicine will be held in Ann Arbor Mich. October 14 through 18 1973. The program will include a full day of educational lectures and video tape on all disciplines in medicine.

For further information regarding this meeting please contact Dr. Michael A. Wainstock, Towsley Center, University of Michigan Medical Center, Ann Arbor, Mich. 48104.

Postgraduate course in angiography

The second postgraduate course in Advances in Angiography Technical Equipment Clinical—Phase II will be held January 30 through February 1 1974 at the Sands Hotel Las Vegas Nevada. The course is sponsored by the Departments of Radiology of Shadyside Hospital in conjunction with the University of Pittsburgh School of Medicine. The course is intended for radiologists, cardiologists and other personnel involved in vascular examinations. Panel discussions by manufacturers, leading investigators will be correlated with clinical comments by prominent clinicians actively involved in vascular examinations. The course will emphasize personal opinions regarding all aspects of arteriography. Registration fee = \$90 (Registration is limited to 500 on a first enrollment basis). Persons wishing further information may write to Shadyside Hospital, Radiology Seminar Department of Radiology, 5230 Centre Ave., Pittsburgh Pa. 15232.

Symposium on diagnosis and management of hypertensive patients

The University of Texas Health Science Center at Houston Medical School and Division of Continuing Education will present a brief review of new developments in the diagnosis and management of the hypertensive patient on November 29 1973. The symposium will be held in the Auditorium of the Jesse H. Jones Library Building at Houston Texas.

Recognizing that important developments have occurred recently to help the physician diagnose and treat the hypertensive patient, this program has been organized. It is designed to help the physician properly apply this new information and also to extend logically information concerning antihypertensive drug therapy to those who have a reasonable chance of benefiting from it. The faculty draws from across the country and addresses itself to these problems and offer acceptable solutions to the practicing physician.

Program director for the symposium is Walter W. Kirlendall, M.D., Professor and Director, Program in Internal Medicine, The University of Texas Health Science Center at Houston Medical School. The registration fee is \$15 including luncheon and handout material.

For further information regarding this program, please write: Office of the Director, The University of Texas Health Science Center at Houston, Division of Continuing Education, P.O. Box 20367, Houston, Texas 77025.

Editorial

Cyclophosphamide and the treatment of the nephrotic syndrome in adults

C S Ogg MD, MRCP

J S Cameron MD FRCP

London England

It is more than 20 years since the publication of the first reports on the use of mechlorethamine (nitrogen mustard) in the treatment of patients with glomerulonephritis.^{1,2} However the difficulties associated with its use and the introduction of corticosteroid therapy reduced interest in the alkylating agents for a decade while the limitations and dangers of steroids were appreciated. By this time new drugs such as cyclophosphamide and chlorambucil which were less toxic than nitrogen mustard and which could be given orally had been introduced and in 1963 the first report of the use of cyclophosphamide in the treatment of children with the nephrotic syndrome was published.³ This report dealt largely with patients suffering from minimal change lesions and although cyclophosphamide has since been used in patients with other renal lesions the great majority of evidence about its value in the treatment of patients with the nephrotic syndrome still concerns this group.

Minimal change lesion

Patients suffering from this condition which is also known as lipid nephrosis usually have a pure nephrotic syndrome without hematuria or hypertension. The

great majority are children but something like 30 per cent of adults with the nephrotic syndrome may also show this appearance.⁴ Examination of biopsy material reveals normal or near normal glomeruli without deposition of immune globulins or changes in the glomerular basement membranes. Usually the condition runs an intermittent course with remissions and relapses occurring spontaneously⁵ or in response to treatment with corticosteroids. In order to maintain remission many patients require continuous therapy with steroids in doses which produce severe side effects. This problem is greatest in children in whom suppression of growth is added to the other well known problems of treatment.

A large number of papers have now been published⁶⁻¹¹ which provide ample evidence that cyclophosphamide is able to induce remission of the nephrotic syndrome in children with this condition. It seemed that these remissions lasted longer than those formerly induced with steroids¹⁰ and in 1970 Barratt and Soothill¹¹ confirmed by means of a controlled trial that this was in fact the case. Patients do still relapse however and it is probable that only in about 50 per cent is permanent remission achieved.^{12,13} Even a temporary result is

From the Renal Unit, Guy's Hospital, London, England.

Received for publication Oct. 2, 1972.

Reprint requests to Dr. C. S. Ogg, Renal Physician, Renal Unit, Guy's Hospital, London SE1, England.

however, well worth having at the least the stigmata of steroid toxicity are given time to regress and growth in height is achieved. Although no trial has been reported which compares cyclophosphamide with azathioprine, the poor results obtained using the latter drug¹⁴ suggest that cyclophosphamide is much superior. Cyclophosphamide may either be given alone¹⁰ or during a remission which has been induced and maintained by steroids.^{11,12} Currently we are using an eight week course of treatment in a dosage of 3 mg per kilogram of body weight per day and have no evidence that larger doses or longer courses influence the long term outlook.^{12,13}

There is much less evidence available about the place of cyclophosphamide in the treatment of adults with minimal renal lesions. This is partly because of the relative rarity of the lesion in this group⁴ but also because the toxicity of steroids in adults is often not as great as it is in children. However the evidence that is available suggests that the situation in adults is no different from that in children.^{15,16} Initial response may be anticipated in the vast majority of patients but again relapse may occur eventually.¹⁶

Occasional patients with biopsies showing minimal renal lesions fail to respond to treatment with steroids. A proportion of these do respond to cyclophosphamide¹⁵ but the majority do not. Many of these patients are probably suffering from a different condition, focal glomerulosclerosis which affects initially only the juxta-medullary glomeruli which may not be present in a biopsy containing only renal cortex. Response of patients with this condition to cyclophosphamide is unusual^{17,18} although not unheard of.¹⁹

Systemic lupus erythematosus

Work on two animal models of this condition, namely the F1 hybrid of the NZB and NZW mouse^{20,21} and Aleutian disease of mink²² in which the animals normally develop a severe proliferative glomerulonephritis has shown that treatment with cyclophosphamide can either slow the progression of the renal disease or prevent its appearance completely. This has provided a rational basis for the use of this drug in the treatment of the nephrotic syndrome

associated with systemic lupus erythematosus in man. Initial uncontrolled reports^{2,24} gave promising results and recently a controlled trial²⁵ has produced clear-cut evidence that cyclophosphamide does have a place in the treatment of these patients. The ideal dose regimen has not yet been established and it is not yet clear that cyclophosphamide is superior to for instance azathioprine and steroids given in combination.

The drug may also have a place in the treatment of other systemic disorders with renal involvement such as Wegener's granulomatosis²⁶ but we have been unimpressed with its efficacy in the treatment of the nephrotic syndrome associated with Schönlein-Henoch purpura.²⁷

Proliferative glomerulonephritis

The majority of adults with the nephrotic syndrome suffer from one or other of the many subtypes of this condition. A common classification, which has been outlined by Cameron²⁸ includes endothelial proliferative nephritis, rapidly progressive nephritis with crescents, and mesangiocapillary glomerulonephritis. It is likely that even these subgroups are heterogeneous and it is not surprising that there is no clear-cut information about the role of cyclophosphamide in the treatment of these conditions. Most patients with the nephrotic syndrome due to proliferative nephritis run a downhill course to renal failure over a period of 10 to 20 years¹² and it would be naive to expect relatively short courses of treatment to achieve more than arrest or slowing of progressive renal functional impairment. However the success of treatment with cyclophosphamide in patients with systemic lupus erythematosus who may show any of these lesions in their renal biopsies provides a continual stimulus towards evaluation of this drug in patients with proliferative nephritis.

We²⁹ and others,³⁰ have treated isolated patients with mesangiocapillary glomerulonephritis with variable but generally unimpressive results while Kincaid Smith and colleagues³¹ have compared patients treated with cyclophosphamide and anti-coagulants together with patients receiving no treatment and showed that the treated group fared better. Patients with biopsies

showing multiple crescents fail likewise to respond to cyclophosphamide alone¹¹ but recently we have used this drug in combination with anticoagulants and steroids with encouraging results.¹²

Membranous nephropathy

The incidence of this lesion among adults with the nephrotic syndrome is variable but in the United Kingdom it accounts for roughly 12 per cent of the cases.⁴ Again patients run a steady downhill course to renal failure over a period of about ten years.¹³ There have been very few reports of the use of cyclophosphamide in this condition but recently Donadio¹⁴ has discussed the result of a small controlled trial which failed to demonstrate any benefit from treatment with this drug. Again however the suggestion that this condition may well have its origin in an immunological disorder as shown by its production in experimental animals¹⁵ and by its occasional appearance in patients with systemic lupus erythematosus should encourage continued interest in therapy with immunosuppressive agents.

Toxicity of cyclophosphamide

The dangers of therapy with cyclophosphamide should not be underestimated. Formerly it was believed that most of the toxicity was reversible and avoidable by careful supervision of the patient with regular checks on the blood count. Nonetheless many patients suffered nausea and alopecia and some died from overwhelming sepsis.¹⁶ Recently however it has been appreciated that permanent damage may be produced to the bladder and to the gonads. Chemical cystitis is a well recognized acute side effect of treatment but it has now been realized that the drug can cause fibrosis of the bladder wall¹⁷ and possibly also bladder tumors.¹⁸ More serious however is the suggestion that cyclophosphamide may cause permanent sterility. Earlier this year Fairley and colleagues¹⁹ reported the results of analysis of seminal fluid from 31 men who had received cyclophosphamide. All showed low or absent sperm counts and testicular biopsy in five showed absent or minimal spermatogenesis. Similar results were obtained by Kumar and colleagues²⁰ from

Belfast and there seems little doubt that cyclophosphamide can cause profound and possibly permanent damage to the adult male testis. The danger of treatment given before puberty is less certain. Berry²¹ was unable to demonstrate any major effect on the subsequent reproductive function of male rats treated with cyclophosphamide before sexual maturation. For obvious reasons there is a paucity of information in boys and in particular there is very little histological information. One boy with Hodgkin's disease who had radiotherapy in addition to 3 months cyclophosphamide was found to have severe testicular atrophy when he died later.²² However we found (unpublished data) normal testes in one child aged 8 years at death who had received 8 Gm of cyclophosphamide over a 180 day period two years previously and Arnel (personal communication) has collected information on the testes of 15 prepubertal boys treated with cyclophosphamide for malignant disease. All were normal except one who had in addition received radiotherapy for his Hodgkin's disease. However Etteldorf²³ has studied 5 men who had received cyclophosphamide during childhood and showed azoospermia in three who had received this drug 6½, 8 and 12 years before. Two of his patients who 8 and 11 years previously had received short courses similar to those now recommended for the treatment of children with the nephrotic syndrome due to minimal renal lesions had satisfactory sperm counts. West²⁴ has seen three men who had received cyclophosphamide in childhood and subsequently fathered children who were normal and it seems that the occurrence of long term testicular damage is dependent on the total dose employed.

Evidence also exists that cyclophosphamide may damage the ovary. Women receiving the drug for more than six months regularly develop amenorrhoea^{25, 26} which may be associated with a low urinary excretion of estrogen and a high urinary excretion of gonadotrophin²⁷ suggesting that cessation of menstruation is due to ovarian failure. However menstruation usually returns when the drug is withdrawn²⁸ and patients have even become pregnant while receiving the drug.²⁹ Pregnancies have

been terminated in two of our own patients who became pregnant while taking cyclophosphamide. This policy was adopted as much for psychosocial as for medical reasons although there is some evidence that cyclophosphamide might be teratogenic in these circumstances.^{44, 47} We (unpublished data) have seen two normal pregnancies following short courses of cyclophosphamide and this has also been the experience of others.⁴⁸

Conclusion

The recognition of the long term effects of cyclophosphamide on the gonads has prompted some revision of ideas about its place in the treatment of patients with the nephrotic syndrome. The drug is most effective in those patients who do best on corticosteroids; hence it should only be given to patients with minimal lesions who experience severe side effects from corticosteroids. Even then it should only be given in short courses particularly when dealing with male patients. In patients with progressive and potentially lethal renal disease the risk of sterility is probably justified provided that the drug can be shown to be more effective than other less toxic agents.

REFERENCES

- Chasis H, Goldring W and Baldwin D S. Effect of febrile plasma typhoid vaccine and nitrogen mustard on renal manifestations of human glomerulonephritis. *Proc Soc Exp Biol Med* 71:565 1949
- Chasis H, Goldring W and Baldwin D S. Effect of nitrogen mustard on renal manifestations of human glomerulonephritis. *J Clin Invest* 29:804 1950
- Coldbeck J H. Experience with alkylating agents in the treatment of children with the nephrotic syndrome. *Med J Aust* 2:987 1963
- Sharpstone P, Ogg C S and Cameron J S. Nephrotic syndrome due to primary renal disease in adults I. Survey of incidence in South East England. *Br Med J* 11:533 1969
- Black D A K, Rose G and Brewer D B. Controlled trial of prednisolone in adult patients with the nephrotic syndrome. *Br Med J* 11:421 1970
- West C D, Hong R and Holland N H. Effect of cyclophosphamide on lipid nephrosis in the human and on aminonucleoside nephrosis in the rat. *J Pediatr* 60:516 1966
- Grupe W E and Heymann W. Cytotoxic drugs in steroid resistant renal disease. *Am J Dis Child* 112:448 1966
- Etteldorf J N, Roy S, Sunnitt R L, Sweeney M J, Wall H P and Berton W M. Cyclophosphamide in the treatment of idiopathic lipid nephrosis. *J Pediatr* 10:158 1967
- Drummond K N, Hillman M A, Marchessault J H V and Feldman W. Cyclophosphamide in the nephrotic syndrome of childhood. Its use in two groups of patients defined by light microscope and immunopathological findings. *Can Med Assoc J* 93:524 1968
- Moncreiff M W, White R H R, Ogg C S and Cameron J S. Cyclophosphamide therapy in the nephrotic syndrome of childhood. *Br Med J* 1:666 1969
- Barratt T M and Soothill J F. Controlled trial of cyclophosphamide in steroid sensitive relapsing nephrotic syndrome of childhood. *Lancet* 11:479 1970
- Barratt T M, Soothill J F, Cameron J S, Chantler C and Ogg C S. A comparative trial of two weeks and eight weeks cyclophosphamide on steroid sensitive relapsing nephrotic syndrome of childhood. *Arch Dis Child* (In press)
- Cameron J S. Bright's disease today. The pathogenesis and treatment of glomerulonephritis. *Br Med J* 1V:87 160 217 1972
- Abramowicz M, Arneil G C, Barnett H C, Barron B A, Edelmann C M, Gordillo G, Greifer I, Hallman N, Kobayashi K and Tidwell H A. Controlled trial of azathioprine in children with the nephrotic syndrome. *Lancet* 1:959 1970
- Udall P R, Feest T G, Morley A R, Tonlinson B E and Kerr D N S. Cyclophosphamide therapy in adults with minimal change nephrotic syndrome. *Lancet* 1:1250 1972
- Cameron J S and Ogg C S. Cyclophosphamide in adult onset nephrotic syndrome. *Lancet* 11:397 1972
- White R H R and Glasgow E. Focal glomerulosclerosis. In Kincaid Smith P, Mathew T and Becker E L, editors. *Glomerulonephritis*. New York, 1972. John Wiley & Sons Inc. (In press)
- Cameron J S, Ogg C S, Turner D R and Weller R O. Focal glomerulosclerosis. In Kincaid Smith P, Mathew T and Becker E L, editors. *Glomerulonephritis*. New York, 1972. John Wiley & Sons Inc. (In press)
- West C D. International Workshop in Immunosuppressive Therapy. Greifer I and Barnett H L, editors. New York, 1972. (In preparation)
- Russell P J, Hicks J D and Burnett F M. Cyclophosphamide treatment of kidney disease in (NZBxNZW) F1 mice. *Lancet* 11:1279 1966
- Russell P J and Hicks J D. Cyclophosphamide treatment of renal disease in (NZBxNZW) F1 hybrid mice. *Lancet* 1:440 1968
- Cheema A, Henson J H and Borham J R. Aleutian disease of mink. *Am J Pathol* 66:543 1972
- Seah C S, Wong K H, Chew A G K and Jayaratnam F J. Cyclophosphamide in the treatment of systemic lupus erythematosus. *Br Med J* 1:333 1966
- Cameron J S, Boulton Jones M, Robinson M O and Ogg C S. Treatment of lupus

- nephritis with cyclophosphamide *Lancet* ii:846 1970
- 25 Steinberg A D Kaltreider H B Staples P J Goetzl E J Talal N and Decker J L Lupus nephritis. A controlled trial *Ann Intern Med* 74:165 1971
- 26 Novack S N and Pearson C M Cyclophosphamide therapy in Wegener's granulomatosis *N Engl J Med* 284:938 1971
- 27 Meadow S R Glasgow E F White R H R Moncrieff M W Cameron J S and Ogg C S Schönlein-Henoch nephritis *Q J Med* 41:241 1972
- 28 Cameron J S Glomerulonephritis *Br Med J* iv 285 1970
- 29 Cameron J S Glasgow E F Ogg C S and White R H R Membranoproliferative glomerulonephritis and persistent hypocomplementaemia *Br Med J* iv:7 1970
- 30 West C D Holland N H McConville J M and McAdams A J Immunosuppressive therapy in persistent hypocomplementaemic glomerulonephritis and in lupus nephritis *Pediatr* 67:1113 1965
- 31 Kincaid Smith P et al *in* Kincaid Smith P Mathew T and Becker E L editors *Glomerulonephritis* New York 1972 John Wiley & Sons Inc. (In press)
- 32 Cameron J S and Ogg C S Rapidly progressive nephritis with extensive crescents *in* Kincaid Smith P Mathew T and Becker E L editors *Glomerulonephritis* New York 1972 John Wiley & Sons Inc. (In press)
- 33 Cameron J S Ogg C S Sargent B and Robertson M S (In preparation)
- 34 Donadio J V International Workshop in Immunosuppressive Therapy Greifer I and Barnett H L editors New York 1972 (In preparation)
- 35 Dixon F J The pathogenesis of immunologically induced nephritis *in* Heptinstall R H ed *Third International Congress of Nephrology* Washington vol 2 Basel and New York 1967 S Karger AG p 97
- 36 Meadow S R Weller R O and Archibald R W R Fatal systemic measles in a child receiving cyclophosphamide for nephrotic syndrome *Lancet* ii 876 1969
- 37 Johnson W W and Meadows D C Urinary bladder fibrosis and telangiectasia after cyclophosphamide therapy *N Engl J Med* 284 290 1971
- 38 Worth P J L Cyclophosphamide and the bladder *Br Med J* iii:182 1972
- 39 Fairley K J Barne J U and Johnson W Sterility and testicular atrophy related to cyclophosphamide therapy *Lancet* i:568 1972
- 40 Kumar R Biggart J D McEvoy J and McGowan M G Cyclophosphamide and reproductive function *Lancet* i:1212 1972
- 41 Berry C L Cyclophosphamide therapy in prepubertal rats and subsequent reproductive performance *Arch Dis Child* 46:709 1971
- 42 Hyman L R and Gilbert E F Testicular atrophy in a prepubescent male after cyclophosphamide therapy *Lancet* ii 426 1972
- 43 Etteldorf J N International Workshop in Immunosuppressive Therapy Greifer I and Barnett H L editors New York 1972 (In preparation)
- 44 Uldall P R Kerr D N S and Tacchi D Sterility and cyclophosphamide *Lancet* i:693 1972
- 45 Cameron J S and Ogg C S Sterility and cyclophosphamide *Lancet* ii:1174 1972
- 46 Greenberg L H Verdes P and Tanaka A R Congenital anomalies probably produced by cyclophosphamide *JAMA* 183 423 1964
- 47 Toledo T M Harper W C and Moser R H Fetal effects during cyclophosphamide and irradiation therapy *Ann Intern Med* 74:187 1971
- 48 Travis L B International Workshop in Immunosuppressive Therapy Greifer I and Barnett H L editors New York 1972 (In preparation)

Early systolic notch in the apexcardiogram in mitral stenosis

Lewis C Becker, MD*

Andrew P Klaus, MD**

J O Neal Humphries, MD

Baltimore, Md

An early systolic notch on the upstroke of the apexcardiogram (ACG) has been considered a useful diagnostic sign of mobile left atrial myxoma.^{1,2} The notch is thought to be caused by movement of the tumor from left ventricle to left atrium in early systole.² Although a small notch in the ACG may sometimes be seen in normal subjects,^{3,4} it has not been reported to occur in mitral valve stenosis.

During evaluation of a patient (K. E.) with auscultatory signs of mitral stenosis an ACG was recorded with an early systolic notch. However an echocardiogram demonstrated the typical findings of mitral valve stenosis and no myxoma; this was confirmed at surgery. Subsequently three additional patients have been found with valvular mitral stenosis in whom an early systolic notch has been recorded.

Methods

Apexcardiograms were obtained by manually holding a 25 mm funnel applicator attached by a 6 cm length of plastic tubing to an air filled Statham 23Db strain gauge on the apical impulse. The patient was most

often in the left lateral position but some times was supine. The signal was amplified by a Hewlett Packard 350 100 C pre amplifier and recorded at 75 mm per second on a Hewlett Packard 4560 Photographic recorder. The damped natural frequency of this system was 8.3 cycles per second determined by a balloon rupture technique.⁵ Simultaneous electrocardiograms (Lead II) and phonocardiograms (lower frequency cut off of 200 Hz) were also recorded.

The preoperative apexcardiograms of all patients who had undergone mitral commissurotomy for relief of mitral stenosis in 1970 through 1971 were reviewed in a retrospective fashion. In these patients no special attempt had been made to record a notch. Of 23 patients who underwent mitral commissurotomy in 1970 through 1971 16 had ACG's and 12 were technically adequate.

In addition the next three patients with pure mitral stenosis proved by catheterization seen in our institution had ACG's in which a definite effort was made to manipulate the funnel over the cardiac apex to

* From the Cardiovascular Division, Department of Medicine, Johns Hopkins University School of Medicine, Baltimore.

Publication Nov 16 1972

to J O Neal Humphries, MD, Cardiovascular Division, Johns Hopkins Hospital, Baltimore, Md.

Resident, Cardiology Division, University of North Carolina School of Medicine, Chapel Hill, N.C.

Dr. Frederick C. Smith, Clin. C, Marion, Ohio

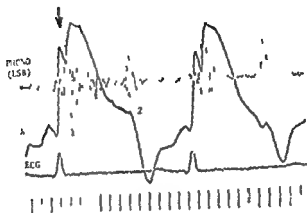


Fig 1 Simultaneous apexcardiogram (ACG) left sternal border phonocardiogram (PHONO) and Lead II electrocardiogram (ECG) are shown for patient E E. The first (1) and second (2) heart sounds are labelled. The prominent notch interrupting the systolic upstroke of the ACG is indicated by an arrow. Time lines at the bottom of the figure are 0.04 second apart.

produce an early systolic notch. The ACG was technically satisfactory in two and unsatisfactory in one. Our original patient (K E) was considered in the prospective group because myxoma was being actively considered at the time of study.

Results

An early systolic notch was seen in two of the twelve adequate ACC's in the retrospective group and in two of the three adequate ACG's in the prospective group. Table 1 presents the clinical data on these four patients and their ACG's are shown in Figs 1 to 4. A prominent notch on the upstroke of the ACC is seen in each of these records. In three patients (E E, K E and R P) the notch coincides with the first heart sound. In the other patient (E S) the notch follows the onset of the first heart sound by 0.03 sec.

The way the pickup was held on the cardiac apex seemed to affect significantly the shape of the ACG and the presence of the early systolic notch. Fig 4 illustrates the ACG obtained in patient K E with the funnel held on the apex in two slightly different positions.

Discussion

Previous reports dealing with the ACC in mitral stenosis^{8,10} have been concerned with the diastolic portion of the recording or with the timing of sounds in the cardiac

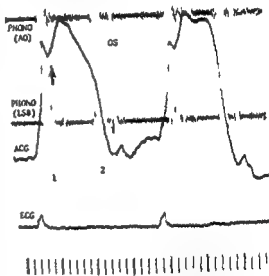


Fig 2 Patient E S PHONO (AO) indicates phonocardiogram in aortic area and OS indicates opening snap. Other abbreviations as in Fig 1.

cycle. However, one of these published ACG's demonstrates early systolic notching (Ref 9 Fig 5).

There are probably several causes of an early systolic notch in the ACG. In some cases the notch may merely represent an artifact of recording. The variability of the systolic portion of the ACG has been previously noted^{8,11} and others have pointed out that marked alterations in the shape of

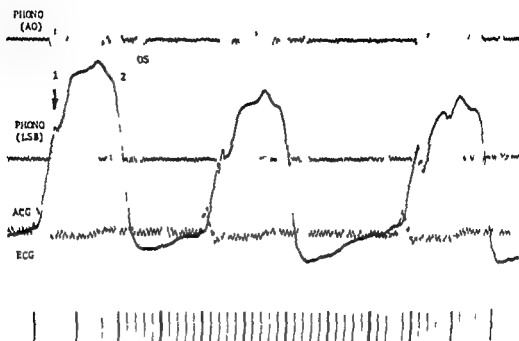


Fig 3 Patient R P For abbreviations see Figs 1 and 2 Note that the patient is in atrial fibrillation and that the early systolic notch is variable In a longer record there was no relation between the cycle length and the prominence of the notch

Table I Clinical data on 4 patients

| Patient | Age (yr) | Sex | MMG | Associated disease | Cardiac rhythm | How discovered | Catheterization† | Surgery† | Q-S ₁ ‡ |
|---------|----------|-----|-----|---------------------|----------------|----------------|------------------|----------|--------------------|
| E E | 49 | F | 12 | Mild AR | NSR | Retropective | + | + | 08 |
| E S | 56 | F | 12 | Mild AR | AF | Retropective | + | + | 08 |
| K E | 25 | F | — | Pregnant | NSR | Propective | — | + | 10 |
| R P | 53 | M | 8 | Mild AS moderate AR | AF | Prospective | + | — | 09 |

* Abbreviations: MMG = mean mitral valve gradient (mm Hg) at cardiac catheterization; AR = aortic regurgitation; AS = aortic stenosis; NSR = normal sinus rhythm; AF = atrial fibrillation.

† Catheterization and Surgery columns indicate how atrial myxoma was excluded in each patient. All patients had mitral valve surgery only.

‡ Q-S₁ = interval in seconds between the onset of the Q wave in the Lead II LCG and the first high frequency component of the first heart sound in the phonocardiogram. Values of Q-S₁ for normals or patients without mitral valve disease are less than 0.06 sec.^{14,15}

the systolic waves may occur with slight changes in the position of the recording device.^{14,15} One of our patients (Fig 4) illustrates that an early systolic notch may be brought out by careful manipulation of the funnel applicator.

In patients with valvular mitral stenosis the notch may reflect the loud first heart sound. It has been shown that there is frequently a prolongation of the interval from the Q wave of the ECG to the onset of the first heart sound in mitral stenosis.^{14,15} This prolonged 'Q-S₁ interval' is thought to be

due to the greater time required for left ventricular pressure to rise above left atrial pressure. Because of the delayed onset of S₁ its low frequency vibrations would occur later on the systolic upstroke of the ACG than normally and might produce a more noticeable distortion.

Our data supports this latter hypothesis since all our patients had prolonged Q-S₁ intervals and increased intensity of the first sound. The notches on the ACG coincided with the S₁ in three of the patients while in the fourth patient the notch followed the

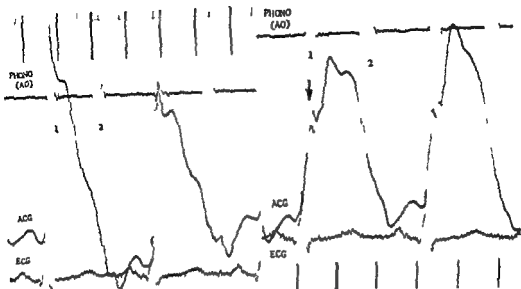


Fig. 4 Patient 4. ■ For abbreviations see Figs 1 and 2. Left panel shows ACG taken in routine fashion. Right panel demonstrates the appearance of a prominent notch after slight movement of the pickup funnel over the cardiac apex.

onset of S_1 by 0.05 second. We have no explanation for this discrepancy in the fourth patient.

Although an early systolic notch in the ACG in patients with clinical sign of mitral valve obstruction has been shown to be non-specific we feel that it should still alert one to the possibility of myxoma and that further investigation should be performed. Ultrasound may be used to examine the echoes from the mitral valve apparatus or echoes from a myxoma tumor mass of the left atrium.^{16,17} If an early systolic notch in the ACG is found in a patient in whom myxoma is a possibility, transseptal puncture of the left atrium should not be done without first visualizing the left atrium during the left phase of a pulmonary arteriogram.

Summary

An early systolic notch on the upstroke of the apexcardiogram (ACG) has previously been reported to occur in mobile left atrial myxoma but has not been noted in valvular mitral stenosis. It has therefore been considered a useful diagnostic sign for myxoma. We report four patients with mitral valve stenosis who had an early systolic notch in the ACG. Myxoma was

excluded by catheterization and/or surgery. We feel that the notch is not specific for myxoma and that its finding should be interpreted with caution.

REFERENCES

1. Zitrnik R S, Giuliana E R and Burchell H B. Left atrial myxoma: phonocardiographic clues to diagnosis. *Am J Cardiol* 23:588 1969.
2. Becker L C and Conti C R. Left atrial myxoma: evidence of tumor movement by apexcardiogram. *Chest* 60:80 1971.
3. Nasser W K, Darr R H, Dillon J C, Tavel M E, Helmen C H, Feigenbaum H and Fisch C. Atrial myxoma II. Phonocardiographic, echocardiographic, hemodynamic and angiographic features in nine cases. *Am Heart J* 83:810 1972.
4. Hahiba K, Ono A, Matsuo S, Yoshioka M, Mori K, Fujiwara C, Yano K and Oshibuchi R. Apexcardiogram and phonocardiogram in the diagnosis of left atrial myxoma: report of a case. *Jap Heart J* 11:202 1970.
5. Mauoch C I, Abbott J A and Rapaport E. Left atrial myxoma with bacteremia: Report of a case with a bifid systolic apical impulse. *Am J Cardiol* 2:133 1970.
6. Coussed V and Epstein E J. The apexcardiogram: its normal features explained by those found in heart disease. *Br Heart J* 25:697 1963.
7. Tafur E, Cohen L S and Levine H D. The normal apexcardiogram: its temporal relationship to electrical, acoustic and mechanical cardiac events. *Circulation* 30:581 1964.

- 8 Fry D L Physiologic recording by modern instruments with particular reference to pressure recording *Physiol Rev* 40:753 1960
- 9 Benchimol A Dimond L G Waxman D and Shen Y Diastolic movements of the precordium in mitral stenosis and regurgitation *Am HEART J* 60:417 1960
- 10 Oreshkov V I A new mechanocardiographic index in evaluation of the severity of mitral stenosis in apexcardiographic study *Am HEART J* 79 789 1970
- 11 Tavel M E Campbell R W Leikenbrium H and Steinmetz L J The apexcardiogram and its relationship to hemodynamic events within the left heart *Br Heart J* 27 879 1965
- 12 Tavel M E Clinical phonocardiography and external pulse recording, Chicago 1972 Year Book Medical Publishers Inc
- 13 Benchimol A Dimond E G and Carson, J C The value of the apexcardiogram as a reference tracing in phonocardiography *Am HEART J* 61 485 1961
- 14 Wells B The assessment of mitral stenosis by phonocardiography *Br Heart J* 16:261 1954
- 15 Kelly J J Diagnostic value of phonocardiography in mitral stenosis mode of production of first heart sound *Am J Med* 19 862 1955
- 16 Kostis J B and Moghadam A A Echocardiographic diagnosis of left atrial myxoma *Chest* 58 550 1970
- 17 Finegan M B and Harrison D C Diagnosis of left atrial myxoma by echocardiography *Am Engl J Med* 282 1072 1970

An analysis of deaths occurring in association with coronary arteriography

Timothy Takaro M D
Herbert N Hultgren M D
David Littmann M D
Elizabeth C Wright
Oleen N C

In the course of a continuing Veterans Administration Cooperative Study of Surgery for Coronary Arterial Occlusive Disease a system of reporting on coronary arteriograms performed by 13 cooperating hospitals since June 1968 was established. These arteriography logs also carried data concerning deaths occurring in association with the procedure. The opportunity thus presented itself of analyzing these data both for incidence and cause of fatalities by reviewing the records and coronary arteriograms of all patients who were reported to us as having died within 10 days of the angiographic procedure. It seemed important to do this because heretofore coronary arteriography has often been reported as being associated with a very low risk.¹⁻⁵ Recent reports in the medical literature and our own data do not bear this out.⁶⁻⁸ We also hoped that preventive measures might be developed if some of the mechanisms of death appeared to be avoidable.

Not all of the 13 participating hospitals were involved in the Cooperative Study for the entire duration of the time span encompassed by this report (June 1968 through June 1972). In addition three

other hospitals not participating in the Study contributed cases. Since the arteriography logs did not require information regarding the technique of coronary arteriography which was employed this was sought retrospectively by questionnaire from each participating hospital; these data are therefore approximations.

Methods

The clinical records, coronary arteriograms, catheterization data, and autopsy reports for all patients reported to have died were reviewed and analyzed. All available coronary arteriograms were personally reviewed. Histopathologic sections of coronary arteries of selected patients were also reviewed. Clinical and autopsy data were tabulated and an attempt was made to correlate the results.

Results

Sixty-six deaths were available for analysis from 16 of the 18 cooperating hospitals. Sixty-four of these were reported from a total of 3,044 arteriograms performed by 17 of these hospitals between June 1, 1968 and June 30, 1972. The incidence of fatalities occurring within 10 days

From the Veterans Administration Hospital, Olean, N. C.

Received for publication Dec. 21, 1972.

Reprint requests to Dr. Timothy Takaro, Veterans Administration Hospital, Olean, N. C. 28860.

of coronary arteriography in these hospitals was thus 2.1 per cent.

All of the 66 patients who died were men, 62 of them between 40 and 60 years of age. Four were over 60. There were 56 white, 9 black, and one native Indian patients. The indication for coronary arteriography was angina pectoris in a patient who was considered a potential candidate for surgery for this symptom in 44 cases. In most instances, angina was of many months' duration. Uncertainty of diagnosis in a patient with atypical chest pain was the indication for arteriography in six, congestive failure or cardiomegaly in four, and postoperative follow-up studies were the indications in eight. In four instances the precise indication was not clear from the record. Only five patients had no angina. There was a history of myocardial infarction in 41 patients (60 per cent) and electrocardiographic evidence of old infarct myocardial ischemia or a positive stress test in 52 (78 per cent).

Left ventriculography was reported as having been performed in 42 patients; there was some abnormality of size, contractility, or ejection fraction in 24 of these (57 per cent). Significant coronary occlusive lesions (narrowing of lumen diameter of 50 per cent or greater) were noted (either on angiograms or at autopsy or both) in all but two patients in this series. In 11 patients the left main coronary artery was significantly stenosed at angiography, and in 31 of 47 in whom coronary angiography had been completed, three or more major vessels were involved with significant lesions. Thus most of the patients who died had significant diffuse disease, and the majority also had subnormal left ventricular function.

Forty-nine patients (74 per cent) died within 24 hours of the completion of the procedure—26 of them (39 per cent) within three hours. We included in this report all deaths reported to us which occurred within 10 days of coronary arteriography.

The exact incidence of fatalities in association with coronary arteriography was difficult to determine for several reasons: (1) arteriography logs were not kept by two hospitals, (2) data on the techniques used in the 3,044 arteriograms were obtained

retrospectively and were imprecise, and (3) there were 13 deaths in patients not registered on the arteriography logs. This occurred because only the names of those patients who underwent coronary arteriography in connection with the Cooperative Study were required to be entered on the logs.

However, precise data about the techniques used in the fatal cases was available in every instance. Based upon these, there were two deaths* (0.3 per cent) in approximately 750 arteriograms using the transbrachial approach, and 51 deaths (2.2 per cent) in approximately 2,300 arteriograms using the transfemoral approach†.

An attempt was made to classify all deaths in accordance with the best available information.

The largest group of deaths (Group 1) (35/66 or 53 per cent) was attributable to acute coronary occlusion, based either on angiographic evidence of sudden occlusion of a previously patent coronary artery (Fig. 1A and B) or pathologic evidence of a fresh occlusive lesion in a major coronary artery (Fig. 2A and B) or both. Of this group, 28/33 or 80 per cent of the patients died within six hours, and five within two days; the remaining two patients died at intervals of three and nine days. In 12 of these patients, an acute myocardial infarction was noted at autopsy, in addition to the occlusive lesion. In 15 of these patients, an embolus with a unique histological appearance was identified.¹³ In all but four instances, the left coronary artery was being catheterized at the time of the acute episode.

Eight patients (Group 2), all of whom died within 24 hours of angiography, were classified as having had an acute cardiac arrhythmia. In none of these patients was an acute myocardial infarction or an acutely occluding coronary arterial lesion identified.

In six patients (Group 3) an acute myocardial infarction was found at autopsy without evidence of an acutely occluding

In an earlier report three deaths were recorded with the transbrachial technique.¹¹ One of these was from a non-V.A. Hospital; the arteriographic volume of this hospital was not used to calculate incidence, and this death is therefore omitted from this calculation.

*p < .001



Fig 1 A and B. A Complete occlusion of the left main coronary artery by embolus (arrow). Moments before during left ventriculography this vessel was seen to be patent. B Complete occlusion of distal right coronary artery by embolus (arrow). On a previous injection of this vessel a few moments earlier the distal right coronary was seen to be patent.

Table 1 Incidence of fatalities in association with coronary arteriography

| | | Number of patients | Per cent deaths |
|---|---|--------------------|-----------------|
| 1 | Ross and Gorlin ⁴ Am Heart Assoc study (80% Cleveland Clinic) 1968 | 3 312 | 0.1 |
| 2 | Waparian and Lehman ⁴ 1967 | 800 | 0.3 |
| 3 | Waltenbach and Lichtlen ¹⁰ 1971 | 1 198 | 0.6 |
| 4 | Selzer Anderson March ⁹ 1971 | 2 275 | 0.9 |
| | San Francisco area | 800 | 2.6 |
| | Another Western city | 891 | 1.0 |
| 5 | Chahine Herman Gorlin ⁹ 1977 | 46 900 | 0.44 |
| 6 | Adams Fraser Abrams ¹¹ 1972 | 3 044 | 2.1 |
| 7 | V.A. Coop Study (up to 10 day) 1972 | | |

arterial lesion either at angiography or at autopsy. Most of these patients (5/6) lived for 12 hours or more (up to 10 days) after the completion of the procedure.

Four patients (Group 4) died of the consequences of a cerebrovascular accident which occurred during the procedure. These patients lived from two to nine days after angiography. Four (Group 5) died in association with problems relating to the femoral arterial puncture, three of them less than 24 hours after the angiographic procedure. One patient (Group 6) died four days after angiography of an acute septicaemia and in eight instances (Group 7) no specific mechanism for death could be identified.

All but one of the 23 patients in Groups

1 to 4 had had at least one change of catheters prior to the acute episode and many had two, three or more changes of catheters. In three instances the fatal episode occurred after change of catheters but prior to the injection of any contrast medium. In seven patients the fatal episode occurred after only a small test injection in or near the orifice of the left coronary artery. In many instances prior to the fatal episode episodes of hypotension or concomitant or associated studies had delayed or prolonged the procedure.

An analysis of more complete data available from a subset of 1297 patients from arteriography logs to determine the risk of arteriography in those with significant stenotic lesions of the left main coronary



Fig 2 A and B A Autopsy specimen showing cut left main coronary artery near its orifice. The lumen is completely occluded by fresh thrombus B Histologic section showing friable coiled platelet material found in this coronary artery (Hematoxylin and eosin Original magnification X25)

Table II Classification of deaths (66 cases)

| | No | Per cent |
|-----------------------------------|----|----------|
| 1 Acute coronary occlusion | 35 | 56 |
| 2 Acute arrhythmia | 8 | 12 |
| 3 Acute myocardial infarction | 6 | 9 |
| 4 Cerebral vascular accident | 4 | 6 |
| 5 Problems related to arteriotomy | 4 | 6 |
| 6 Acute septicemia | 1 | 1.5 |
| 7 Non specific or unidentified | 8 | 12 |

Table III Acute coronary occlusion in association with coronary arteriography

| | No |
|--|------------|
| Fresh thrombosis described at autopsy (catheter embolus identified) | 26 (15) |
| Acute hemorrhage into atheromatous plaque | 2 |
| Acute coronary arterial subintimal dissection | 1 |
| Angiographic evidence only | 6 |

artery was possible. In this subset there were 125 patients with significant lesions of the left main—eight of whom died in association with arteriography (6.4 per cent). This compares with 26 deaths from among the remaining 1,170 patients with out left main coronary artery lesions (2.2 per cent) ($p < 0.01$).

Discussion

The incidence of fatal complications was so low earlier using the transbrachial tech-

nique that coronary arteriography quickly gained acceptance as a relatively safe diagnostic procedure.¹⁻¹⁴ However, more recently there has been an unmistakable increase in the incidence of fatal complications which seems to be related to increasing use of the transfemoral approach.⁶⁻¹¹

The disparity between the fatal complication rates of the two techniques has been noted by several authors.^{7-10,14} For example Chalune and associates⁷ reported no deaths in 413 studies using a brachial arteriotomy and a 2 per cent incidence of fatal complications among 478 studies using the transfemoral technique. There is little doubt that the transfemoral method has been easier to teach and to learn since it utilized the familiar Seldinger technique for gaining access to the femoral artery and preformed easily guided catheters were used.¹⁵ The complications involving brachial arteriotomy which can be subintimal even in good hands could be avoided by using the transfemoral technique.¹⁵ It seemed to be merely a simple extension of the very useful technique of selective catheterization of the branches of the abdominal aorta.

Unfortunately, in the ascending aorta, and with multiple changes of catheters the problem is more complex and the procedure can be more dangerous. The explanation for the disparity between the fatal complication rates following transbrachial and transfemoral coronary arteriography seems to be related to the mechanism of sudden coronary arterial occlusion due to

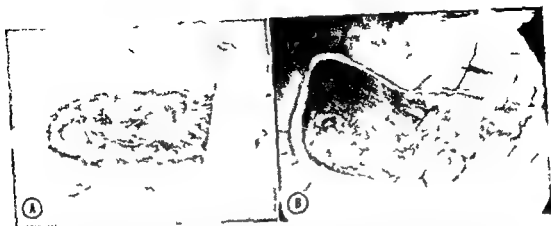


Fig 3 A and B Examples of coronary arterial emboli from two patients who died in association with coronary arteriography. Note characteristic appearance of coiled masses of platelets with few red cells and little fibrin in an artery with little evidence of severe atherosclerosis (Hematoxylin and eosin. Original magnification $\times 25$).

Table IV Influence of technique on mortality

| | Transbrachial | | Transfemoral | |
|---|---------------|-----------|---------------|-----------|
| | % of patients | Incidence | % of patients | Incidence |
| Kaltenbach and Lichtlen ¹ (1971) | 1 367 | 0.4% | 431 | 0.9% |
| Chahine-Herman-Gorlin ² (1972) | 413 | 0.0% | 478 | 2.0% |
| VA Coop Study (1972) | 750 | 0.3% | 2 300 | 2.2% |

emboli from guidewires or catheters. As will be noted the guidewire is the essential element in this mechanism and guidewires are not ordinarily used in the transbrachial technique.

In a previous report¹² 26 patients were reported who had died in association with coronary arteriography and in whom evidence of acute coronary occlusion was found. Catheter embolism was the postulated mechanism in many instances. In a subsequent report¹³ it was shown that catheter emboli could be identified by their unique and characteristic histologic appearance which was completely unlike that seen with *in situ* thrombi found in some cases of sudden death from coronary arterial disease.

Since our initial report nine additional cases of acute coronary occlusion apparently due to this mechanism have been contributed by participating hospitals. The

characteristic histologic appearance of these emboli is attributed to the way they are thought to have developed (Fig 3A and B). It is known that fibrin and platelets begin to adhere to the surfaces of catheters and presumably of guidewires (if the transfemoral technique is being employed) very soon after these foreign objects are exposed to the blood stream.^{14,20} In removing a guidewire from a catheter this fine layer of platelets is easily stripped from the wire by the closely fitting catheter tip and may adhere to the outside of the catheter tip as an accumulation undetectable by pressure monitoring or flushing. This platelet material may then be deposited in or swept into a coronary artery (or a brachiocephalic vessel) when the catheter tip nears one of these orifices (Fig 4).^{12,18} In addition the fine sleeve of platelets which is deposited on the outside surface of a catheter is wiped off this surface by the arterial wall at the

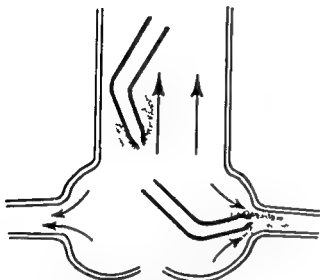


Fig 4 Diagrammatic representation of one mechanism postulated for the propagation of catheter emboli: Thrombotic material wiped off guidewire adheres to the outside surface of the tip of the catheter. The catheter tip (upper black heavy lines) is now in the ascending aorta with the adherent thrombotic material streaming in the direction of blood flow (vertical arrows). Embolization from this position can result in a cerebral vascular accident. In the lower right the catheter tip is near a coronary arterial orifice. Blood flow (curved arrows) now causes thrombotic material to stream toward the coronary arteries and to cause coronary embolization even without any injection of contrast material. (Reproduced from Takaro et al: Acute coronary occlusion following coronary arteriography: mechanisms and surgical relief. *Surgery* 72:1018, 1972. The C V Mosby Co.)

arterial puncture site and accumulates at this site as the catheter is withdrawn (Fig 5A, B, C and D). There is considerable radiologic evidence to support this hypothesis.²⁰ The thrombotic material adheres to the injured arterial wall and if a guide wire has been replaced in the artery in preparation for the introduction of a second catheter, it adheres also to the retained guidewire. The tip of the second catheter may sweep this material off the guidewire as it is introduced into the artery. In most instances, this material is almost certainly swept away by the blood stream, fragmenting and causing undetectable microscopic peripheral emboli. In rare instances it adheres to the tip of the second catheter and causes either cerebral or coronary arterial embolization. It appears in the lumen of the coronary vessels as a mass of material consisting of tightly packed coils or cords

of aggregated platelets without significant fibrin or red cells.¹¹ We believe that two of the cases reported by Giddings and associates² represent emboli of this nature. Such an appearance was also noted in two surviving cases in which the embolus was surgically extracted within hours of the coronary arteriographic procedure.¹²

Further angiographic evidence to substantiate the possibility of adherence of a sleeve of platelets to a catheter tip is provided by the unusual circumstance of the pocketing of contrast medium in such a sleeve, rendering it radiopaque (Fig 6A, B, and C). Less obvious angiographic evidence of the adherence of thrombotic material to a catheter tip is not infrequently noted by the alert angiographer. It appears sometimes as a fleck of contrast material adhering to the outside of the catheter, or as a small area of radiolucency in a bolus of contrast medium. It is clearly distinguishable from an air bubble by its irregular configuration. In one instance the progress of this radiolucent material down a coronary artery and its subsequent lodgment was recorded on an angiogram. In this instance it resulted in a non fatal infarction.

A third postulated mechanism for the development and propagation of catheter emboli is shown diagrammatically in Fig 7, involving the so called pig tail catheters used for left ventriculography. Long, stringy, radiopaque thrombotic material adhering to the pigtail catheter was recorded on cineangiography in at least one instance. There were no untoward sequelae.

In most instances angiographic evidence of coronary embolization consisted of the abrupt appearance during the procedure of total occlusion of a coronary artery shown to be patent only moments before or of stasis or retention of contrast material in a coronary artery after the rest of the material had disappeared (Fig 1). Some of these cases have been assumed to represent subintimal deposition of contrast material, but without pathologic proof.^{21,22} We have been able to uncover only one pathologically proved case of demonstrable subintimal dissection of the coronary artery during coronary arteriography with deposition of contrast medium in the arterial

wall and conclude that most such appearances represent instead coronary arterial embolization.²⁸

We believe that the guidewire is an essential element in the pathogenesis of catheter embolism (Figs 4, 5 and 7). It is necessary for the transfer of thrombotic material from a foreign body surface (either wire or catheter) to the tip of a cardiac catheter. For this reason it is difficult to postulate such a mechanism as occurring using the transbrachial technique with an open brachial arteriotomy and without a guidewire.

The probability of increased platelet deposition is increased by prolongation of a study either because of technical difficulty, lack of experience or for special studies involving ventricular function or implant or graft opacification. It is also increased by stasis of blood flow which would be expected to occur in patients either with hypotension or poor ventricular function and low cardiac output or in vessels involved with atherosclerosis or with both.^{27, 28} Some of the patients in Groups 2 and 3 may also have had coronary emboli from catheters, but the resuscitative efforts may have fragmented such emboli so that they were not recognized or pathologic examination may have been incomplete. Certainly significant hemodynamic effects²⁹ as well as serious arrhythmias due to the contrast medium itself³⁰ or to the irritation of catheters or to reduction of coronary blood flow by catheter tips which partially or completely occlude coronary arterial orifices have been recognized but the relationship between these mechanisms and the deaths in our patients in Groups 2 and 3 must remain conjectural.

It is noteworthy that so many patients in this series had diffuse disease in two or three of the major branches of the coronary artery and a substantial proportion had evidence of significant left ventricular dysfunction. The highest risk appeared to be in patients with significant stenoses of the left main coronary artery, a fact previously noted by Cohen and colleagues.³¹ They reported a 16 per cent incidence of fatality in association with coronary arteriography in a small group of patients with similar lesions of the left main coronary artery.



Fig 5 A through D Diagrammatic representation of another mechanism postulated for the formation and propagation of catheter emboli. A Catheter (heavy black lines) passing through wall of artery (fine double lines) and lying in arterial lumen. The intra arterial portion of the catheter is covered by a fine deposit of platelets here exaggeratedly thick (stippled layer). B A guidewire has been inserted into the lumen of the catheter which is now being withdrawn from the artery. The layer of platelets is being wiped off the catheter by the arterial wall and is accumulating at the arterial puncture site. C The thrombotic platelet material remains adherent to the puncture site and around the retained guidewire. D This material is being wiped off the guidewire by the tip of the next catheter inserted into the artery. This material adheres now to the tip of this second catheter. (Reproduced from Takaro et al. Acute coronary occlusion following coronary arteriography: mechanisms and surgical relief. *Surgery* 72:1018, 1972. The C. V. Mosby Co.)

Hildner and colleagues³² in a provocative paper entitled "Pseudocomplications of cardiac catheterization" reported 26 acute events including 12 sudden deaths in patients who were scheduled for cardiac catheterization but did not have the procedure. These acute problems included acute myocardial infarctions, acute arrhythmias, cerebral vascular accidents, peripheral arterial occlusion, etc. strikingly similar types of complications to those which occurred in their catheterized patients following the procedure and also to our series. They concluded that one should not presume technical error or operator negligence as the cause of all complications following a cardiac catheterization procedure unless such is demonstrable. In any large group of patients it is likely that chance complications will occur as a result of the patient's disease rather than as the result of the procedure. This may well have been true for some of the deaths reported



Fig 6 A through C Sequential frames from cineangiogram showing Judkins type coronary arterial catheter tip in descending thoracic aorta an unusually long sleeve of clot (arrows) fortuitously opacified by contrast material is shown clinging to the catheter tip and waving in the blood stream This can easily become an embolus

in our series. However, in those cases where an acute coronary occlusion not previously present but demonstrated to have occurred during the angiographic procedure was identified, or a coronary embolus was seen at autopsy an iatrogenic cause for the death must be ascribed.

Since our earlier report¹² a number of additional case reports of emergency surgery for acute coronary occlusion or acute myocardial ischemia occurring during coronary arteriography have appeared in the medical literature.²⁶⁻²⁷ Survival was usually reported in patients who were not in cardiogenic shock at the time of surgery. There was usually a fatal outcome for those in shock. In almost all instances the transfemoral technique had been used. Cheng⁸ writes that, from his personal conversations with many coronary arteriographers, fatal thromboembolism following selective coronary arteriography is much more frequent than has been generally acknowledged. In one of his cases (as well as in one of the series reported by us) the coronary arteries were essentially normal at autopsy—except for the embolic occlusion of a coronary artery.

Prevention of fatal complications

Since the incidence of fatal complications as reported here and elsewhere is so much greater following the transfemoral technique, use of the transbrachial technique may prevent some of these deaths. If the transfemoral technique is used shortening the procedure minimizing associated studies⁸ and avoiding the use of a pigtail type catheter in left ventriculography prior to selective catheterization of the coronary arteries, and the use of non thrombogenic catheters or coatings²⁸ may be additional safeguards. Avoiding the use of a guide wire by using a Teflon sheath to re enter the femoral artery allowing free bleeding between changes of catheters to permit the washing away of withdrawal thrombi at the femoral arterial puncture site or removal of the guidewire and use of the opposite femoral artery for change of catheters or for ventriculography have all been suggested.²⁹ The use of systemic heparin during the procedure has also been advocated⁴⁰⁻⁴¹ however while blood clotting due to stasis of blood in catheters may thus be prevented heparin does not alter platelet aggregation nor presumably, platelet deposition on

catheter surfaces.^{13, 14} Therefore one can not be certain that systemic heparinization will surely prevent acute coronary occlusion from catheter emboli. Indeed in one of our patients fatal coronary embolization occurred in spite of the administration of 50 mg of heparin intra arterially. It would appear that a combination of heparin plus an agent which would retard platelet aggregation on foreign surfaces would be most effective if the transfemoral technique is used.

Treatment

The surgical treatment of acute coronary occlusion occurring in association with coronary arteriography has been discussed.¹⁵ Emergency embolectomy and aortocoronary saphenous vein bypass graft insertion was followed by survival of a number of patients, mostly those without cardiogenic shock at the time of the surgery. Since emergency surgery may be necessary following coronary arteriography it is probably unwise to perform this procedure regularly in large volumes in a hospital not equipped for emergency cardiac surgery using extracorporeal circulation.

Total cost of coronary arterial surgery

If the estimate of 35 000 aortocoronary bypass grafts performed since the technique was introduced five years ago is correct, it seems likely¹⁶ then it is also very probable that over 100 000 coronary arteriograms have been performed to identify these cases. If an estimate of 1 or 2 per cent as the average risk of fatality with coronary arteriography is also correct and this is also quite probable then approximately 1 000 to 2 000 patients have succumbed in association with this diagnostic test. The problem is obviously of some considerable magnitude.

Coronary arteriography is undoubtedly an invaluable diagnostic tool which provides morphologic information obtainable in no other way. It is worth noting that this information is obtained at some cost, however. This increases the total mortality rate for which surgery must be held accountable since most of these coronary arteriograms would not have been performed if surgery were not being offered as possible



Fig. 7 Diagrammatic representation of a third mechanism postulated for the formation and propagation of catheter emboli. Platelet thrombus in area of stagnant blood flow in tip of pigtail catheter used for ventriculography may also adhere to the guidewire used for replacing this catheter with a coronary arterial catheter and may be thus transferred to the tip of the second catheter. Embolization may occur as noted in Fig. 5 (Reproduced from Takaro et al. Acute coronary occlusion following coronary arteriography: mechanisms and surgical relief. *Surgery* 72:1018, 1972. The C. V. Mosby Co.)

therapy. For this reason also fatal complications of coronary arteriography while they probably cannot be completely eliminated, must be kept to an absolute minimum.

Summary and conclusions

In a Cooperative Study involving 18 hospitals over a four year period the incidence of fatalities occurring in association with coronary arteriography (up to 10 days following the procedure) was 2.1 per cent. It was 0.3 per cent using the transbrachial technique and 2.2 per cent using the transfemoral.

This disparity is explained by the mechanism of coronary or cerebral arterial embolization from platelet deposits on guide wires and catheters for which evidence is presented in a majority of cases. Other mechanisms causing death were acute cardiac arrhythmias, acute myocardial infarction, complications of the femoral arterial puncture, septicemia and unknown causes. Patients with significant stenoses of the left main coronary artery were at the greatest risk (6.4 per cent). Prevention of such complications depends upon the technique used, speed, skill and possibly the use of systemic anticoagulant therapy or antithrombogenic catheters and guide wires.

Coronary arteriography should not be undertaken lightly since the risk of a fatal complication at least using the trans

femoral technique, is appreciable. This fact is borne out not only by our study, but by reports from a number of other recently reported large series.

The risk of coronary arteriography should be reckoned as part of the 'total cost' of coronary arterial surgery.

Arteriography logs and/or cases were contributed for this study by the following VA Hospitals: Ann Arbor, Brooklyn, Buffalo, Dallas, Hines, Lexington, Little Rock, Long Beach, Los Angeles, Madison, Minneapolis, Oteen, Palo Alto, Salt Lake City, San Francisco, Seattle, and Washington, and also by Loyola University Hospital. The cooperation of the cardiologists, surgeons, angiographers, and pathologists at these Hospitals is gratefully acknowledged. We are also very grateful to R. L. Crislip, M.D., of Portland, Oregon, who loaned the films from which Fig. 6 was taken.

REFERENCES

1. Sones F, Mason C. Cine coronary arteriography: anesthesia and analgesia. 46:199, 1967.
2. Judkins M P. Percutaneous transfemoral selective coronary arteriography. *Radiol Clin North Am* 6:467, 1968.
3. Green G, Gerald S, McKinnon C, Malcolm R, Sch Judkins M. Complications of selective percutaneous transfemoral coronary arteriography and their prevention. *Circulation* 45:552, 1972.
4. Ross R, Richard S, and Gorlin R. Coronary arteriography. *Circulation* 37 and 38 (Suppl III):67, 1968.
5. Takaro T, Dart C H Jr, Scott S M, Fish R G, and Nelson W M. Coronary arteriography—indications, techniques, complications. *Ann Thorac Surg* 21:213, 1968.
6. Abrams H, Herbert L. Coronary arteriography: complications and indications. In: *Questions and Answers*. JAMA 219:917, 1972.
7. Chahine R A, Herman M V, and Gorlin R. Complications of coronary arteriography: comparison of the brachial to the femoral approach. (Abstract) *Ann Intern Med* 76:862, 1972.
8. Cheng T O. Fatal thromboembolism following selective coronary arteriography. (Editorial) *Chest* 62:1, 1972.
9. Selzer A, Anderson W I, and March H L. Indications for coronary arteriography—risks vs. benefits. *Calif Med* 115:1, 71.
10. Kaltenbach M, and Lichtlen P. *Coronary heart disease*. Stuttgart, Germany, 1971. Georg Thieme Verlag, pp 19, 29.
11. Wechsler A S, and Sabiston D C. Coronary bypass grafts for myocardial ischemia. *Franz Ingelfinger ed. In: Controversy in internal medicine II* (In press).
12. Takaro T, Pifarre R, Wuerflein M D, Hall A D, Gage A A, Scott S M, Dart C H Jr, and Price H P. Acute coronary occlusion following coronary arteriography: mechanisms and surgical relief. *Surgery* 72:1018, 1972.
13. Price H P, Sollod N, Scott S M, and Takaro T. Unusual coronary emboli associated with coronary arteriography. *Chest* (In press).
14. Kasparian H, and Lehman J S. Coronary arteriography in myocardial infarction. *Radiol Clin North Am* 3:453, 1967.
15. Campion B C, Frye R, Robert L, Pluth J R, Furbairn J F, and Davis G D. Arterial complications of retrograde brachial arterial catheterization. *Mayo Clin Proc* 46:589, 1971.
16. Formanek G, Gustave F, Robert S, and Amplatz K. Arterial thrombus formation during clinical percutaneous catheterization. *Circulation* 41:833, 1970.
17. Glancy J, John J, Fishbone G, Heinz M, Ralph N. Nonthrombotic arterial catheters. *Am J Roentgenol* 108:716, 1970.
18. Jacobsson B, Bergentz S, Erik S, and Ljungqvist U. Platelet adhesion and thrombus formation on vascular catheters in dogs. *Acta Radiol (Diagn)* 8:221, 1969.
19. Nachmani G H, Lessin L S, Motomiya T, and Jensen W N. Scanning electron microscopy of thrombogenesis on vascular catheter surfaces. *N Engl J Med* 286:139, 1972.
20. Nejad M S, Klapar M A, Steggerda F R, and Gianturco C. Clotting on the outer surfaces of vascular catheters. *Radiology* 91:248, 1968.
21. Jacobsson B, Paulin S, and Schlossman D. Thromboembolism of leg following percutaneous catheterization of femoral artery for angiography—symptoms and signs. *Acta Radiol (Diagn)* 8:97, 1969.
22. Siegelman S S, Caplan L H, and Annes G. Complications of catheter angiography—study with Ocillometry and Pullout angiograms. *Radiology* 91:251, 1968.
23. Giddings J A, See J R, Lewis R D, and Cosby R S. Thromboembolism following coronary arteriography. *Chest* 61:235, 1972.
24. Braunwald E. Deaths related to retrograde arterial catheterization or coronary arteriography, or both. *Circulation* 37 and 38 (Suppl III):23, 1968.
25. Hias J M, Peterson C R, and Jones R C. Subintimal dissection of the coronary arteries. *Circulation* 38:678, 1968.
26. Pifarre R, Spinazzola A, Nemickas R, Scanlon P J, and Tobin J R. Emergency aortocoronary bypass for acute myocardial infarction. *Arch Surg* 103:525, 1971.
27. Jacobsson B, Paulin S, and Schlossman D. Thromboembolism of leg following percutaneous catheterization of femoral artery for angiography. *Acta Radiol* 8:109, 1969.
28. Lang E, and K. A survey of complications of percutaneous retrograde arteriography. *Seldinger technique*. *Radiology* 81:257, 1963.
29. Friesinger G C, Schaffer J, Criley J M, Gaertner R A, and Ross M S. Hemodynamic consequences of the injection of radiopaque material. *Circulation* 31:730, 1965.
30. Snyder C F, Formanek A, Frech R S

- and Amplatz K. The role of sodium in promoting ventricular arrhythmia during selective coronary arteriography. *Am J Roentgenol* 113:567 1971
- 31 Cohen M V, Cohn P F, Herman M V and Gorlin R. Diagnosis and prognosis of main left coronary artery obstruction. *Circulation* 45 and 46 (Suppl 1) 57 1972
- 32 Hildner F J, Javier R P, Ramaswamy K and Samet P. Pseudocomplications of cardiac catheterization. *Chest* (In press)
- 33 Berger R L, Wong J and Messer J V. Coronary embolectomy for acute embolization. *Am J Surg* 123:726 1972
- 34 Cohn L H, Fogarty T J, Daily P D and Shumway N E. Emergency coronary artery bypass. *Surgery* 10:821 1971
- 35 Cohn L H, Gorlin R, Herman M V and Collins J J. Aorto-coronary bypass for acute coronary occlusion. *J Thor Cardiovasc Surg* 64:503 1972
- 36 Reul G J, Morris G C Jr, Howell J F, Crawford E S and Stelter W J. Current concepts in coronary artery surgery (A critical analysis of 1287 patients). *Ann Thorac Surg* 14:243 1972
- 37 Scanlon P F. Myocardial revascularization during acute phase of myocardial infarction. *JAMA* 218:207 1971
- 38 Amplatz K. Simple nontrombogenic coating. *Invest. Radiol* 6:280 1971
- 39 Fisher Vincent and Shah A. Safe performance of coronary arteriography by retrograde femoral (Judkins) technique. Personal communication
- 40 Nelson R M and Osborne A G. Systemic heparinization for percutaneous catheter arteriography (Abstract). *Circulation* 43 and 44 (Suppl 11) 205 1971
- 41 Wallace Sidney, Medelin H, De Jongh D and Gianturco C. Systemic heparinization for angiography. *Am J Roentgenol* 116:704 1972
- 42 Lessin L S, Jensen W N, Kelsner G A, Motomiyama T, Nachmani G H. Vascular catheters and thrombogenesis. *N Engl J Med* 287:468 1972
- 43 Salzman E W. The limitations of heparin therapy after arterial grafting operation. *Surgery* 57:131 1965
- 44 Adams D F, Fraser D and Abrams D. Hazards of coronary arteriography. *Seminars in Roentgenology* 11:357 1972

Left ventricular gallop sound and acute myocardial infarction

Charles P. Riley, MD

Richard O. Russell, Jr., MD

Charles E. Rackley, MD

Birmingham, Ala

Pulmonary rales and ventricular gallop sound (VGS) and cardiogenic shock have important prognostic implications in patients with acute myocardial infarction. The classification of patients with acute myocardial infarction into four groups on the basis of these bedside findings, as proposed by Killip and Kimball¹ has proved to be quite useful in predicting immediate survival. Class IV patients, those with severe left ventricular power failure, have an extremely poor prognosis. Intensive studies are now being made of the possible benefits of volume expansion,² circulatory assistance,^{3,4} or vein bypass surgery^{5,7} during the acute phase in these patients. The purpose of this study was to correlate a ventricular gallop sound heard in patients with acute myocardial infarction at the time of admission to the Coronary Care Unit with survival during the acute phase. The clinical objective was to identify a group of patients who might benefit from more intensive therapeutic efforts.

Methods

During a 24 month period 156 patients with proved myocardial infarction were admitted to the Myocardial Infarction Re-

search Unit at the University of Alabama in Birmingham. Criteria for diagnosis included a compatible clinical history with the development of diagnostic Q waves or evolution of ST segment and T wave abnormalities and characteristic changes in SGOT and LDH activity in serial blood samples. Patients with acute myocardial infarction were divided into three groups on the basis of their initial electrocardiogram.^{8,9} Anterior myocardial infarction included patients with electrocardiographic evidence of either acute anteroapical or anterolateral infarction. Inferior myocardial infarction included patients with either diaphragmatic or posterior QRS changes. Patients in the third group with ST changes only were felt to have had definite myocardial necrosis on clinical grounds with a typical rise and fall in serum enzymes (SGOT and/or LDH), but without electrocardiographic evidence of transmural infarction. Patients who presented with left bundle branch block or a nonspecific intraventricular conduction defect were included in this last group as well as patients with subendocardial infarction (myocardial necrosis).

Patients were classified on bedside exami-

From the Myocardial Infarction Research Unit, Cardiovascular Research and Training Center, University of Alabama Medical Center.

Supported by the Myocardial Infarction Research Unit Contract PH 43-67-1441 and the Cardiovascular Research and Training Center Program Project Grant HE 11-310, National Institutes of Health; Department of Health, Education and Welfare.

Received for publication Jan. 2, 1973.

Reprint requests to Charles E. Rackley, MD, Department of Medicine, University of Alabama Medical Center, University Station, Birmingham, Ala. 35294.

nation according to the myocardial infarction classification of Killip and Kimball.¹ An uncomplicated patient (Class I) had neither pulmonary rales nor a ventricular filling gallop sound. Patients with mild or moderate congestive heart failure (Class II) exhibited basilar pulmonary rales persistent after coughing and/or a ventricular filling gallop sound detected by at least one of the investigators. Pulmonary edema (Class III) was defined as the presence of moist rales over more than half of the lung fields. Shock associated with myocardial infarction (Class IV) was diagnosed clinically when there was evidence of severely decreased peripheral perfusion such as cool clammy skin, weak pulse, mental obtundation or oliguria and a systolic blood pressure below 90 mm Hg or 80 mm Hg below previous blood pressure.

A ventricular gallop sound was considered present if it was audible by at least two observers on the day of admission to the Coronary Care Unit. Phonocardiograms were not recorded on these patients and therefore the presence or absence of a ventricular gallop sound is based on clinical observations. The accuracy of the senior authors in the recognition of a ventricular gallop sound has been confirmed by comparison to phonocardiography in previous studies.¹⁰

Pulmonary artery pressure was measured in 49 patients admitted with acute myocardial infarction. In 4 additional patients a catheter was introduced into the brachial artery and advanced to the left ventricle for measurement of left ventricular end diastolic pressure. After explanation of the procedure to the patient and/or relatives informed consent was obtained. Cardiac catheterization was performed primarily in patients with clinical heart failure and/or shock and the detection of a VGS alone or with pulmonary rales was considered clinical evidence of heart failure. The patient remained in a radiolucent bed and a ceiling mounted fluoroscope was used during catheter manipulation. Under sterile conditions and local anesthesia a cutdown was made on an antecubital vein. Early in the investigation standard radiopaque cardiac catheters were used but later were replaced by Swan Ganz catheters. In the right atrium 50 mg of lidocaine was injected and the catheter was advanced to the

Table 1 Ventricular gallop sound (VGS) in relationship to infarct location and mortality rate

| | | Survived | Died | Total |
|-------------|-----|----------|------|-------|
| VGS present | ANT | 22 | 16 | 38 |
| | INF | 7 | 2 | 9 |
| | ST | 9 | 7 | 16 |
| VGS absent | ANT | 31 | 8 | 39 |
| | INF | 31 | 4 | 35 |
| | ST | 17 | 2 | 19 |
| | | 117 | 39 | 156 |

Abb: ST as ANT = anterior myocardial infarction INF = inferior myocardial infarction ST = myocardial necrosis

pulmonary artery. The mid position of the interposterior chest thickness was used as reference for the level of the right atrium.

A chi square test of independence was performed for each of the variables shown in the tables comparing mortality rate with the presence or absence of a ventricular gallop sound. A t test was performed to determine the significance of different mean left ventricular filling pressure in patients with or without ventricular gallop sounds.

Results

A ventricular gallop sound was audible in 63 of 156 patients (40 per cent) at the time of admission to the Coronary Care Unit (Table 1).

The pulmonary artery or left ventricular end-diastolic pressure was measured in 53 patients with acute myocardial infarction (Fig 1). In 27 patients a VGS was audible and in 26 the sound was not detected. The time delay from admission to the Unit and measurement of pulmonary artery pressure ranged from 1 to 84 hours but 42 patients (79 per cent) were studied within 24 hours. There was no significant difference in delay of study in patients with a VGS (13.5 ± 20.7 hours) or those without the finding (18.6 ± 19.1 hours). In the 26 patients without a VGS the pulmonary artery end diastolic pressure (PAEDP) or left ventricular end diastolic pressure (LVEDP) ranged from 2 to 44 mm Hg and was less than 12 mm Hg in 11 patients. In the 27 patients with a VGS 25 revealed a PAEDP or LVEDP > 12 mm Hg and 2 possessed a PAEDP within the normal range 7 and

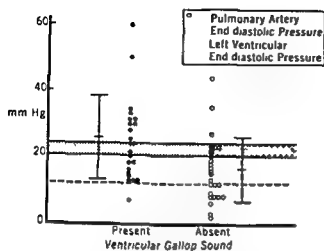


Fig 1 Ventricular gallop sound and left ventricular filling pressure in acute myocardial infarction (dashed line = 12 mm Hg shaded bar = 20 to 24 mm Hg) Twenty five of 27 patients with a VGS demonstrated abnormal elevations of PAEDP or LVEDP whereas 11 of 26 patients without a VGS revealed normal left ventricular filling pressure. Patients with a VGS and PAEDP or LVEDP < 20 mm Hg could have received a volume expander for improvement of left ventricular function and patients with a VGS and PAEDP or LVEDP > 24 mm Hg would be candidates for blood volume reduction for optimal ventricular performance.

11 mm Hg respectively. In the 25 patients with a VGS and an abnormally elevated PAEDP or LVEDP above 12 mm Hg, 8 patients revealed PAEDP or LVEDP < 20 mm Hg and 13 patients demonstrated a PAEDP or LVEDP > 24 mm Hg. An audible VGS in patients with acute myocardial infarction was associated with an abnormally elevated PAEDP or LVEDP in 25 of 27 patients. Mean left ventricular filling pressure was higher in those with a VGS (25.1 ± 12.6) than in those without a VGS (16.3 ± 9.65) ($P < .005$).

In hospital deaths occurred in 40 per cent of patients with a ventricular gallop sound, as opposed to 15 per cent of patients without a ventricular gallop sound ($P < .001$). A ventricular gallop sound was present in 14 of 20 patients (70 per cent) with cardiogenic shock (Table II). Four of six patients (67 per cent) in cardiogenic shock without a ventricular gallop sound and 11 of 14 (79 per cent) with this physical finding died in the hospital ($P > .05$). Although a left ventricular gallop sound was associated with an increased risk of death in the total group of patients studied, this physical finding was not significantly associated with death from cardiogenic shock.

A twofold increase in mortality rate was

noted in patients with anterior myocardial infarction and a ventricular gallop sound ($P < .05$). A fourfold increase in mortality rate was noted in patients with ST changes only—i.e., myocardial necrosis, and a ventricular gallop sound ($P < .05$) (Table I). Of the seven patients who died with myocardial necrosis and a ventricular gallop sound, only three died in cardiogenic shock. A ventricular gallop sound was heard on admission in 11 of 15 patients who subsequently died with left ventricular power failure. Therefore, a left ventricular gallop sound present on admission to the hospital was not useful in predicting which patients would subsequently die in cardiogenic shock.

The presence of a ventricular gallop sound in patients with anterior and ST segment infarctions was associated with an unfavorable prognosis for survival. There was no significant association between a ventricular gallop sound and death due to cardiogenic shock ($P > .05$). The increased risk of death in patients with anterior or ST segment infarction is largely attributable to deaths due to causes other than cardiogenic shock. Only patients with inferior infarction seemed to be at increased risk of death from cardiogenic shock in the presence of a ventricular gallop sound. However, the number of patients with inferior infarction was too small for these results to be statistically significant.

Discussion

Although ventricular gallop sounds are useful in the clinical recognition of heart failure, the prognostic importance of this auscultatory finding has not been delineated in patients with acute myocardial infarction. Thus, while a ventricular gallop sound may be associated with an overall increase in mortality rate in patients with acute myocardial infarction, this finding does not have high predictive value for death in the individual patient. It is important to note that patients having ST segment changes only—i.e., myocardial necrosis, in association with a ventricular gallop sound have an increased hospital mortality rate. Thus, increased alertness for arrhythmias or depressed left ventricular function must be maintained in this group of patients who are otherwise generally considered to have an uncomplicated clinical course.

A left ventricular diastolic gallop sound is well recognized as a clinical hallmark of left ventricular failure. The genesis of this sound remains obscure but is generally felt to represent an imbalance between the volume of left ventricular inflow during the rapid filling phase of diastole and the ability of the left ventricle to accommodate the amount of inflow.^{11,12} This sound may result from sudden tension upon the mitral valve apparatus as the ventricle rapidly expands and elongates.¹³

Porter and colleagues¹⁴ have shown that left ventricular wall stress values at peak left ventricular filling rates in patients with chronic valvular heart disease and a ventricular gallop sound may be normal. This supports the concept of a relatively compliant left ventricle in this group of patients although direct measurements of left ventricular compliance remain under investigation.¹⁴ One patient in Porter's series however with acute mitral regurgitation and left ventricular failure had a very high wall stress during peak left ventricular filling. This would suggest a relatively non-compliant left ventricle and perhaps a different mechanism for the ventricular gallop sound. These investigators also found that patients with primary myocardial disease and a ventricular gallop sound had elevated wall stress values during peak LV filling. An abnormal elevation in wall stress during this phase of diastole would indicate a non-compliant left ventricle and thus another mechanism for the ventricular gallop sound than in those patients with relatively compliant left ventricles during early diastole. The frequent occurrence of atrial gallop sounds in patients with acute myocardial infarction is taken as evidence for reduced left ventricular compliance.¹⁵ However the hemodynamics of atrial gallops in acute infarction remain to be delineated and many of these patients also have ventricular gallop sounds.

In the present investigation 25 of the 27 patients with a VGS demonstrated an abnormal PAEDP or LVEDP greater than 12 mm Hg. Ratshin and associates¹⁶ have previously reported in patients developing shock with acute myocardial infarction that a VGS was always associated with a left ventricular or pulmonary artery end diastolic pressure greater than 12 mm Hg.

Table II Ventricular gallop sound (VGS) and mortality rate in patients with cardiogenic shock (n = 20)

| | Survived | Died | Total |
|-------------|----------|------|-------|
| VGS present | 3 | 11 | 14 |
| VGS absent | 2 | 4 | 6 |
| Total | 5 | 15 | 20 |

Russell and co-workers¹⁷ have demonstrated that in patients with acute infarction left ventricular filling pressure can be elevated with blood volume expansion and ventricular performance can be improved. In Russell's studies optimal improvement in left ventricular function occurred with LVEDP or PAEDP in a range of 20 to 24 mm Hg. In the present study in the 25 patients with a VGS and PAEDP or LVEDP > 12 mm Hg 8 patients revealed PAEDP or LVEDP < 20 mm Hg and 13 patients exhibited a PAEDP or LVEDP > 24 mm Hg. The patients with a VGS and PAEDP or LVEDP < 20 mm Hg could have received a volume expander in order to improve left ventricular function and the patients with a VGS and PAEDP or LVEDP > 24 mm Hg would be candidates for blood volume reduction for optimal ventricular performance. Additional pharmacologic observations emphasize the need for accurately assessing left ventricular function in acute myocardial infarction. Several recent clinical studies have failed to demonstrate significant hemodynamic improvement following digitalis administration in acute myocardial infarction.^{18,19} Experimental and clinical investigations have shown an increased sensitivity to digitalis with toxic arrhythmias during the immediate post infarction period.^{20,21} Therefore a patient with acute myocardial infarction and a VGS should be suspected of having depressed left ventricular function which could be improved with volume expansion or reduction. However measurement of pulmonary artery pressure is necessary to identify these patients and to select the appropriate agents for improving left ventricular function. The present investigation confirms the need for hemodynamic monitoring in patients with a VGS in the presence of mild or severe heart

failure as well as in cardiogenic shock. The best treatment for such patients should be based on initial and continuous measurements of left ventricular hemodynamics.

Summary

A ventricular gallop sound, pulmonary rales, and cardiogenic shock have important prognostic implications in patients with acute myocardial infarction. The purpose of this study was to examine the relationship between VGS and death in myocardial infarction. A VGS was audible in 63 of 156 patients (40 per cent) with proved myocardial infarction admitted to a Myocardial Infarction Research Unit during a 24 month period. Fifty three patients were studied hemodynamically and 25 of 27 patients with an S₃ gallop demonstrated an abnormally elevated pulmonary artery end diastolic pressure greater than 12 mm Hg. Hospital deaths occurred in 40 per cent of patients with a VGS versus 15 per cent of those without this auscultatory finding. Seventy five per cent of patients in cardiogenic shock died but a VGS was not significantly associated with death in this group. Although in all patients a VGS was associated with increased mortality rate deaths were not due to cardiogenic shock. This study confirms the need for hemodynamic monitoring in patients with acute myocardial infarction who exhibit a VGS as evidence of left ventricular power failure. Treatment should be based upon both clinical observations and continued evaluation of the hemodynamic alterations associated with left ventricular power failure.

REFERENCES

- 1 Killip T and Kimball J T Treatment of myocardial infarction in a coronary care unit. *Am J Cardiol* 20:457 1967
- 2 Porter C M, Karp R B, Russell R O Jr and Rackley C E Pulmonary artery monitoring in cardiogenic shock. *Arch Intern Med* 127:304 1971
- 3 Krakauer J S, Rosenbaum A, Freed P S, Jaron D and Kantrowitz A Clinical management of ancillary to phase shift balloon pumping in cardiogenic shock. *Am J Cardiol* 27:123 1971
- 4 Jacoby J A Results of counterpulsation in patients with coronary artery disease. *Am J Cardiol* 27:137 1971
- 5 Anagnostopoulos C E and Kittle C F The surgical aspects of acute myocardial infarction. *Surg Clin North Am* 51:69 1971
- 6 Hill D J, Kerth W J, Kelly J J, Selzer A, Armstrong W, Popper R W, Langston M F and Cohn K E Emergency aortocoronary bypass for impending or extending myocardial infarction. *Circulation* 43 and 44 (Suppl 1):105 1971
- 7 Mundth E D, Yrachak P M, Buckley M J, Leimbach R C, Kantrowitz A and Austen W G Circulatory assistance and emergency direct coronary artery surgery for shock complicating acute myocardial infarction. *N Engl J Med* 282:1382 1970
- 8 Friedberg C K Diseases of the Heart Third ed Philadelphia and London 1966 W B Saunders Company p 817
- 9 Estes E H The Heart Second ed New York 1970 McGraw Hill Book Company p 312
- 10 Porter C M, Baxley W A, Eddleman E E Jr, Frimer M and Rackley C E Left ventricular dimensions and dynamics in patients with gallop heart sounds. *Am J Med* 50:721 1971
- 11 Kuo P T, Schnabel T G, Binkmore W S and Wheat A F Diastolic gallop sounds: the mechanism of production. *J Clin Invest* 36:1035 1957
- 12 Cravasse L, Wheat M W, Wilson J R, Leeds R F and Taylor J W The mechanisms of the generation of the third and fourth heart sounds. *Circulation* 23:635 1962
- 13 Fleming J S Evidence for a mitral valve origin of the left ventricular third heart sound. *Br Heart J* 31:192 1969
- 14 Smith M, Russell R O Jr, Feld B J and Rackley C E Left ventricular compliance and abnormally contracting left ventricular segments following myocardial infarction. *Clin Res* 20:398 1972
- 15 Hill J C, O'Rourke R A, Lewis R O and McGarahan G M The diagnostic value of the atrial gallop in acute myocardial infarction. *Am Heart J* 78:194 1969
- 16 Ratshin R A, Rackley C E and Russell R O Jr Hemodynamic evaluation of left ventricular function in shock complicating myocardial infarction. *Circulation* 45:127 1972
- 17 Russell R O Jr, Rackley C E, Pombo J, Hunt D, Potania C and Dodge H T Effects of increasing left ventricular filling pressure in patients with acute myocardial infarction. *J Clin Invest* 49:1539 1970
- 18 Milmerona R, Schroder G and Werko L Hemodynamic effects of digitalis in acute myocardial infarction. *Acta Med Scand* 180:55 1966
- 19 Balcon R, Hoy J and Sowton E Hemodynamic effects of rapid digitalization following acute myocardial infarction. *Br Heart J* 30:373 1968
- 20 Ratshin R A, Russell R O Jr and Rackley C E Cardiovascular response to digoxin in complicated and uncomplicated myocardial infarction. *Circulation* 44 (Suppl 1):215 1971
- 21 Morris J J, Taft C V, Whalen R E and McIntosh H D Digitalis and experimental myocardial infarction. *Am Heart J* 77:342 1969
- 22 Wolf M J, Scheidt S and Killip T Heart failure complicating acute myocardial infarction. *Circulation* 40:1125 1972

Coronary arteriographic findings in patients with axis shifts or S-T-segment elevations on exercise stress testing

Frederick N Hegge MD

Narp Tuna MD PhD

Howard B Burchell MD PhD

Minneapolis Minn

When transient myocardial ischemia occurs either in spontaneous attacks of angina pectoris or induced by an imposed stress such as exercise or hypoxia it has been long known that there may be electrocardiographic changes expressed mainly as shifts in the ST segment. The usual positive response is a depression in the ST segment in the standard and/or precordial leads.

The description and extent of shift couched in the term depression of the ST segment has been the subject of many papers with focus on its specificity and discriminatory power in the identification of inadequate coronary perfusion and coronary disease. Data concerning the use of exercise tests are voluminous and in the United States one of the most popular exercise tests has been that devised by Masters and others and much information concerning its diagnostic and prognostic value is available.^{1,2} One of us (Burchell³) has reviewed the usefulness of the exercise test in the different approaches of clinician, epidemiologist and work physiologist.

The ST-segment shift related to is

chemia when it occurs is usually perpendicular to the long axis of the heart so that one may conceive of a shell of ischemic myocardium presumably proximate to the endocardium which is relatively more ischemic than that proximate to the epicardium thus creating a mean injury dipole expressed as an ST-segment depression in the standard leads and elevation in the right arm unipolar lead and depression of the left precordial leads. Modifications of the latter leads (e.g. right shoulder to apical area) are commonly used while monitoring the electrocardiographic response during exercise. It has been long noted that occasionally the electrocardiographic response to induced stress could be manifested by ST-segment elevation in the precordial leads or in leads subtending posterior inferior wall of the heart resulting in an electrocardiogram (ECC) simulating acute infarction. Such patterns have also been seen in individuals having spontaneous angina particularly those variants characterized by prolonged pain at rest a phenomenon in patients emphasized as distinctive by Prinzmetal and

From the Section of Cardiology Department of Medicine University of Minnesota, Minneapolis, Minn.
Supported by the National Heart and Lung Institute Grant HL-08527-09

Received for publication Jan. 2 1973

Reprint requests: Narp Tuna, MD University Hospital Box 481 Minneapolis Minn. 55455

Table I *Electrocardiographic (resting and postexercise) and coronary angiographic findings in sub*

| Patient | Age | Sex | Resting ECG | | | S T-segment elevation (mm) | | | | | | | | | |
|---------|-----|-----|------------------------|-----|-----|----------------------------|-----|-----|-----------------|-----------------|-----------------|----------------|----------------|----------------|----------------|
| | | | Normal or minimal ST T | AMI | DVI | I | II | III | aV _R | aV _L | aV _F | V ₁ | V ₂ | V ₃ | V ₄ |
| E A | 47 | M | \ | | | | | 2 | | 1 | | | | | |
| B S | 42 | M | \ | | | | | 0.5 | | 1 | | | | | |
| G B | 49 | M | \ | | | | 0.5 | 1 | | 1.5 | | | | | |
| K C | 44 | M | \ | | | | | | | | | 1 | | | |
| A F | 62 | M | \ | | | | | | | | | 1 | 2 | 2 | 2 |
| V K | 62 | M | \ | | | | | | | | | 3 | 3 | 3 | |
| G L | 60 | M | \ | | | | | | | | | 1.5 | 2.5 | 4 | 1 |
| M W | 66 | F | \ | | | | | | | | | 1.5 | 2 | 1.5 | |
| M M | 37 | M | \ | | | | | | | | | 1.5 | | | |
| L B | 44 | M | | \ | | | | | | | | | | | 2.5 |
| R K | 38 | M | | \ | | | | | | | | 1 | 2 | 2 | 2 |
| L N | 49 | M | | \ | | | | | | | | 1.5 | 2 | 3 | |
| A P | 50 | M | | \ | | | | | | | | 2 | 2 | | |
| J F | 37 | M | | | \ | | | 0.5 | | 0.5 | | 2 | 5 | | |
| C M | 56 | M | | | \ | | | | | | | | | | |
| P H | 45 | M | | \ | \ | | 0.5 | 1 | | 1 | | | | | |
| H N | 60 | M | | \ | \ | | | | | | | 2.5 | 1.5 | 2 | |
| C S | 38 | M | | \ | \ | | 0.5 | 1 | | 0.5 | | 1.5 | 1.5 | 2.5 | 3.5 |

□ A. obstruct = coronary artery obstruction Rt = right LAD = left anterior descending LC = left circumflex

associates¹ and Silverman and Flamm.² The reasons underlying this effect have been conjectured but obviously it suggested an obstructive lesion localized predominantly to one of the major coronary branches. With the availability of coronary arteriograms one has been allowed to make correlations between this unusual ST-segment shift and the locus of the coronary obstruction and this was the basic objective of this study. A similar study has been reported by Fortuin and Friesinger.³

In some instances exercise may produce aberration in the pattern of excitation of the left ventricle, including complete left bundle branch block. There may also be more subtle changes which one can attribute to disturbances in excitation pathways in the proximal or distal portions of the left bundle fasciculus. These alterations are expressed primarily in the change in axis deviation in the frontal plane, and an attempt has been made in this study to relate these changes when they occurred to the pattern of coronary arterial obstruction.

Materials and methods

This study was based on a retrospective review of the findings in 158 patients who had both treadmill tests and coronary arteriograms at the University of Minnesota Hospitals prior to Sept 1 1971. The purpose of the study was to evaluate the association between coronary arteriographic findings and exercise induced ST-segment elevations or axis shifts on maximal treadmill exercise tests.

The treadmill tests done in all 158 patients were multistage, maximal exercise tolerance tests according to Doan and co-workers⁴ in which the ECG was monitored by a single bipolar lead during exercise and by a full 12 lead ECG which was recorded in the supine position on a multichannel machine immediately before and after exercise and at frequent intervals thereafter. All patients who were found to have impressive exercise induced ST-segment elevations or impressive exercise induced left or right axis shifts were included in the study. In the patients with exercise induced ST-segment elevations the height and ECG location of the eleva-

with postexercise ST segment elevation

| | | Coronary artery obstruction | | | Collaterals |
|----------------|----------------|-----------------------------|-----|-----|--------------|
| V ₁ | V ₂ | Rt | LAD | LC | |
| | | 95 | 40 | 60 | To right |
| | | 99 | 90 | 100 | LAD to LC |
| | | 90 | 0 | 0 | None |
| | | 0 | 85 | 0 | None |
| | | 25 | 100 | 0 | Rt to LAD |
| | | 50 | 100 | 90 | None |
| | | 50 | 95 | 75 | None |
| | | 50 | 95 | 0 | Rt to LAD |
| | | 0 | 0 | 0 | None |
| | | 0 | 50 | 0 | None |
| | | 100 | 100 | 45 | Rt to LAD |
| | | 0 | 100 | 0 | Minimal |
| | | 100 | 100 | 25 | Rt to LAD |
| | | 90 | 90 | 90 | None |
| | | 100 | 100 | 100 | To all 3 |
| | | 100 | 100 | 100 | To all 3 |
| | | 100 | 95 | 95 | To Rt |
| 2 | | 100 | 100 | | To Rt to LAD |

tions were tabulated along with the coronary arteriographic findings. In the patients with exercise induced axis shifts the magnitude of the shift was determined on the hexaxial reference system by using the QRS magnitude before and after exercise in Leads I and III and in Leads I and aV_F. The direction and magnitude of the axis shifts were tabulated along with the coronary arteriographic findings.

The coronary arteriograms of 154 of the 158 patients had already been reviewed as part of a larger continuing study. In all 154 patients the degree of obstruction of the right left anterior descending and left circumflex coronary arteries had been graded on a scale of 0 to 4 with 0 being no obstruction, 1 standing for 1 to 24 per cent obstruction, 2 equaling 25 to 49 per cent obstruction, 3 being 50 to 99 per cent obstruction and 4 complete obstruction. The presence and the extent of collateral circulation was also recorded in each patient. This system for grading coronary artery obstruction was used in the tabulation of the location and severity of the coronary artery disease in the groups of

patients with exercise induced axis shifts. However, in the group of patients with exercise induced ST-segment elevations there was an additional review of the coronary arteriographic findings and an estimate of the actual percentage, rather than grade of coronary artery obstruction was included in the tabulation of the location and severity of the coronary artery disease.

Results

Exercise induced ST-segment elevations

Exercise induced ST-segment elevations which were impressive to us were found in 18 of the 158 patients studied (Table 1). These ST-segment elevations were present in the precordial leads only in 12 patients (Fig. 1) in the inferior limb leads only in 5 patients (Fig. 2) and in both the precordial and the inferior limb leads in one patient. The ST-segment elevations ranged from 1 to 5 mm in the precordial leads and from 0.5 to 2 mm in the inferior limb leads (standardization 1 cm = 1 mv). In 9 of the 18 patients there was no evidence of a previous myocardial infarction on the resting ECG. But of the remaining 9 patients 4 had ECG evidence of a previous anterior infarction, 2 had ECG evidence of a previous inferior infarction and 3 had ECG evidence of both a previous anterior and inferior infarction (Table 1).

When the coronary arteriograms of the 13 patients with exercise induced precordial ST-segment elevations were reviewed it was found that there was at least an 80 per cent obstruction of the left anterior descending artery in 11 of the 13 patients (Table II). In addition in a twelfth patient there was 50 per cent diffuse obstruction of the left anterior descending artery along with normal right and left circumflex arteries and evidence of a previous anterior myocardial infarction on the resting ECG.

Of the six patients with precordial ST-segment elevations and no evidence of a myocardial infarction on resting ECG five had at least an 80 per cent obstruction of the left anterior descending artery but none had severe obstruction of the right coronary artery. The sixth patient in this group had normal coronary arteriograms and the least impressive ST-segment elevations of all the patients studied. In this

patient, there was an S T-segment elevation of 1.5 mm, which was present only in Lead V₁ and which was associated with reversal of the T wave direction from inverted to upright (Fig 3)

Of the 7 patients with precordial S T-segment elevations and evidence of a previous myocardial infarction on the resting ECG 6 had at least an 85 per cent obstruction of the left anterior descending artery, and the seventh had a 50 per cent diffuse obstruction as discussed above. But in 5 of these 7 patients there was also severe generalized disease with an 85 per cent obstruction of the right coronary artery as well as the left.

When the coronary arteriograms of the 6 patients with exercise induced inferior limb lead S T-segment elevations were reviewed, it was found that there was at least an 85 per cent obstruction of the right coronary artery in all 6 patients (Table III). This severe obstruction was present only in the right coronary artery in 2 of the 3 patients with no evidence of a previous myocardial infarction on the resting ECG. But there was at least an 85 per cent obstruction in all 3 of the major coronary arteries in each of the 3 patients with evidence of a previous myocardial infarction on the resting ECG.

In 16 of the 18 patients with localized exercise induced S T-segment elevations these changes were associated with at least an 85 per cent obstruction of the compatible coronary artery. In a seventh patient they were associated with a 50 per cent diffuse obstruction of the compatible coronary artery and no obstruction of the other two major vessels. Hence there was severe obstruction of the compatible coronary artery in 17 of the 18 patients with exercise induced S T-segment elevations. And as discussed previously the only patient who did not have severe obstruction of the compatible vessel also had the least impressive exercise induced S T-segment elevations in the entire group.

The severe obstruction of the coronary artery compatible with anterior or inferior ischemic localization was accompanied by varying degrees of obstruction of the other major coronary arteries. In those patients with no evidence for a previous myocardial infarction on the resting ECG severe ob-

struction was often present only in the compatible coronary artery. This was the case in 7 of the 9 patients who had no previous infarction on the ECG. But in those patients who had evidence of a previous myocardial infarction on the resting ECG, severe obstruction of the compatible coronary artery often co-existed with severe obstruction of the other major coronary arteries as well. This was the case in 7 of the 9 patients who had a previous infarction on the ECG.

Exercise induced axis shifts. Exercise induced axis shifts were noted on visual inspection in 13 of the 158 patients studied. In 9 of the patients there was an exercise induced right axis shift (Fig 4) and in 4 of the patients there was an exercise induced left axis shift (Fig 5). In the 9 patients with an exercise induced right axis shift the change in axis with exercise ranged from a change of as little as +14 degrees to a change of as much as +91 degrees (Table IV). In the 4 patients with an exercise induced left axis shift the change in axis after exercise ranged from a change of as little as -11 degrees to a change of as much as -56 degrees (Table V). In addition there was a slight prolongation of the QRS complex after exercise in 2 of the 4 patients with exercise induced left axis shifts.

A tabulation of the coronary arteriographic findings in the 9 patients with exercise induced right axis shifts did not demonstrate any tendency toward predominant localization of coronary artery obstruction in any particular vessel (Table IV). Also as the magnitude of the exercise induced right axis shift increased there was no consistent increase in the severity of the coronary artery obstruction found on arteriography (Table IV). And furthermore the coronary artery obstruction found in these 9 patients was if anything less severe than that found in the total group of patients analyzed. Only 4 of the 9 patients with exercise induced right axis shifts had a 50 per cent or more obstruction of at least one major coronary artery as compared to 103 of the total group of 154 patients analyzed (Table VI).

A tabulation of the coronary arteriographic findings in the 4 patients with exercise induced left axis shifts did demon-

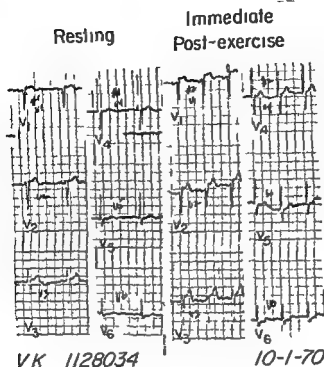


Fig 1 Pre and postexercise ECGs of V K (precordial leads only) showing transient exercise-induced S-T-segment elevations in Leads V₁ to V₆

Table 11 Frequency of at least 85 per cent obstruction in patients with exercise induced precordial ST elevation

| | Resting ECG | | | | Total |
|--------------------------|-------------|---------------------|---------------------|----------------------------------|-------|
| | No infarct | Anterior infarction | Inferior infarction | Anterior and inferior infarction | |
| Number of patients | 6 | 4 | 11 | 2 | 13 |
| Right coronary | 0 | 2 | 1 | 2 | 5 |
| Left anterior descending | 5 | 3 | 1 | 2 | 11 |
| Left circumflex | 1 | 0 | 1 | 2 | 4 |

An additional patient in this group had 50 per cent obstruction of the left anterior descending artery with a normal right and left circumflex arteries.

strate a tendency toward localized coronary artery obstruction (Table V). It was found that 3 of these 4 patients had severe coronary artery disease with complete obstruction of the left anterior descending artery. However, the number of patients in this group was too small for a satisfactory comparison with the larger group (Table VI).

Discussion

Exercise induced ST-segment elevations
Reversible nonexercise induced ST-segment elevations in leads subtending the epicardial surface have previously been reported in association with severe coronary artery disease. In their review of the subject Fortuin and Friesinger⁷ cited the

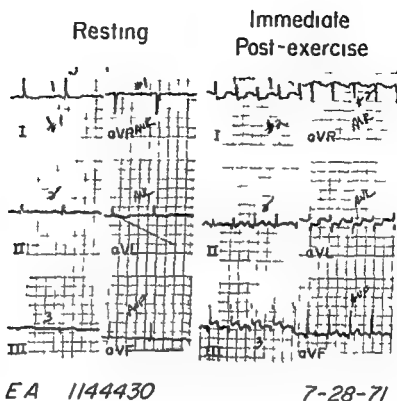


Fig 2 Pre and postexercise ECGs of E A (extremity leads only) showing transient exercise-induced S-T-segment elevations in Lead III and aV_F.

Table III Frequency of at least 80 per cent obstruction in patients with exercise induced inferior S T elevation

| | Resting ECG | | | | |
|--------------------------|---------------|---------------------|---------------------|----------------------------------|-------|
| | No infarction | Inferior infarction | Anterior infarction | Anterior and inferior infarction | Total |
| Number of patients | 3 | 1 | 0 | 2 | 6 |
| Right coronary | 3 | 1 | 0 | 2 | 6 |
| Left anterior descending | 1 | 1 | 0 | 2 | 4 |
| Left circumflex | 1 | 1 | 0 | 2 | 4 |

results of various animal experiments is supporting evidence for the importance of this association. They noted that temporary ligation of a coronary artery has been shown to produce reversible S T-segment elevations in dogs⁹ and that these changes usually do not occur unless the blood flow is reduced by at least 70 per cent.¹⁰ They also noted that in that variant of angina pectoris particularly emphasized by Prinzmetal and colleagues³ reversible S T-segment elevations are associated with

severe coronary artery disease on coronary arteriographic or postmortem examinations.

Reversible S T-segment elevations on exercise ECGs have been reported in association with severe localized coronary artery disease and also in association with ventricular aneurysms. Fortuin and Friesinger reported total or near total obstruction of the compatible coronary artery in 11 of 12 patients with localized exercise induced S T-segment elevations. It is note

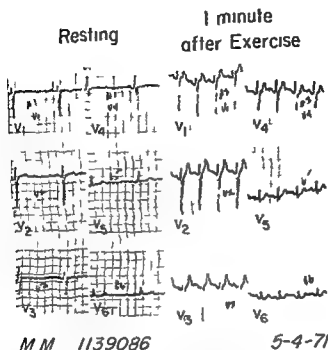


Fig 3 Pre and postexercise ECGs of M M (precordial leads only) showing transient exercise induced S-T-segment elevations in Lead V₂ in association with reversal of the T wave direction from inverted to upright

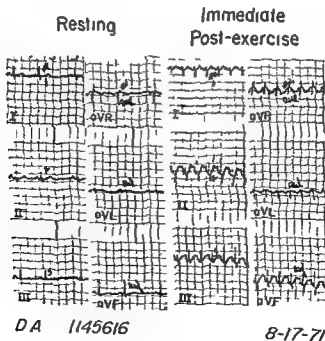


Fig 4 Pre- and postexercise ECGs of D A (extremity leads only) showing a transient exercise induced right axis shift.

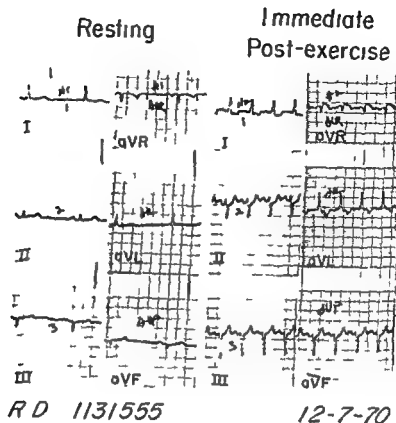


Fig 5 Pre and postexercise ECGs of R D (extremity leads only) showing a transient exercise-induced left axis shift

worthy that only one of these 12 patients had a history consistent with Prinzmetal's variant of angina pectoris. Gorlin and associates¹¹ reported ST-segment elevations in 16 of 24 patients with ventricular aneurysms, and in 4 of the 16 patients with ST-segment elevations these changes were noted only after a Master's two step test. Rubenstein and co-workers¹² reported abnormal left ventriculograms in 8 of 11 patients with ST-segment elevations on exercise ECG. Localized left ventricular dysfunction was found in 8 of the 9 patients with a previous myocardial infarction but this finding was not present in the 2 patients without a previous infarction.

In the University of Minnesota series exercise induced ST-segment elevations were associated with severe obstruction of the coronary artery compatible with the localization of the ischemic zone in 17 of the 18 patients studied. Furthermore the severe obstruction was limited to the compatible coronary artery in 7 of the 9 patients without a previous infarction seen on the ECG, but it was present in both the compatible coronary artery and in

other major coronary arteries as well in 7 of the 9 patients with a previous infarction on ECG. Hence it appears that severe coronary artery disease is likely to be limited to the compatible coronary artery in patients without a previous infarction on the ECG. But it appears that severe coronary artery disease is likely to be present in multiple coronary arteries including the compatible coronary artery, in patients with a previous infarction on ECG. The presence of collaterals connecting with the most diseased artery ("compatible coronary artery") did not seem to alter the exercise induced ST-segment elevation.

Exercise induced axis shifts. Acute transient right and left axis shifts have previously been reported after the injection of radiopaque dye during coronary arteriography¹³ after acute myocardial infarction¹⁴ and after exercise tolerance testing.^{15,16} Both right and left exercise induced axis shifts have been reported in patients presumed to be free of coronary artery disease¹⁵ and in patients known to have angina pectoris.¹⁶

1/1 m 86
A m 5

Table IV Grade of coronary artery obstruction in patients with exercise induced right axis shift

| Patient | Resting ECG | Treadmill test | Axis shift (degrees) | Axis (degrees) | | C A obstruct | | | | Collateral | Postexercise QRS prolongation |
|---------|------------------|----------------|----------------------|----------------|---------------|--------------|----|-----|----|------------|-------------------------------|
| | | | | Pre exercise | Post exercise | Rt | LM | LAD | LC | | |
| F T | ST T changes | + | +14 | +41 to | +55 | 0 | 0 | 0 | 0 | - | - |
| J A | Normal | - | +22 | +61 to | +83 | 0 | 0 | 0 | 0 | - | - |
| F S | LVH ST T changes | + | +25 | +16 to | +41 | 4 | 2 | 3 | 4 | + | - |
| R J | LVH ST T changes | + | +38 | +47 to | +85 | 0 | 0 | 3 | 1 | - | - |
| R E | ST T changes | + | +39 | +35 to | +94 | 0 | 0 | 1 | 0 | - | - |
| H H | Normal | + | +59 | +19 to | +78 | 2 | 0 | 1 | 1 | - | - |
| W E | Normal | + | +68 | +37 to | +105 | 3 | 3 | 3 | 3 | - | - |
| D A | Normal | - | +79 | +62 to | +141 | 0 | 0 | 0 | 0 | - | - |
| B S | Normal | + | +91 | +22 to | +113 | 3 | 0 | 3 | 1 | + | - |

C A obstruct = coronary artery obstructive Rt = right LM = left main LAD = left anterior descending LC = left circumflex
LVH = left ventricular hypertrophy

*0 = No obstruction 1 = 1 to 24 per cent 2 = 25 to 49 per cent 3 = 50 to 99 per cent 4 = complete obstruction.

Table V Grade of coronary artery obstruction in patients with exercise induced left axis shift

| Patient | Resting ECG | Treadmill test | Axis shift (degrees) | Axis (degrees) | | C A obstruct | | | | Collateral | Postexercise QRS prolongation |
|---------|--------------|----------------|----------------------|----------------|---------------|--------------|----|-----|----|------------|-------------------------------|
| | | | | Pre exercise | Post exercise | Rt | LM | LAD | LC | | |
| L N | ASMI | + | -11 | -36 to | -47 | 0 | 1 | 4 | 0 | + | + |
| V C | Normal | - | -31 | -23 to | -54 | 0 | 0 | 0 | 0 | 0 | - |
| R D | ST T changes | + | -36 | -9 to | -45 | 3 | 0 | 4 | 4 | + | + |
| V h | ST T changes | + | -56 | +40 to | -16 | 2 | 0 | 4 | 3 | + | - |

C A obstruct = coronary artery obstructive Rt = right LM = left main LAD = left anterior descending LC = left circumflex
ASMI = anterior septal myocardial infarction

*0 = No obstruction 1 = 1 to 24 per cent 2 = 25 to 49 per cent 3 = 50 to 99 per cent 4 = complete obstruction.

Several aspects of the trifascicular nature of the ventricular conduction system suggest the possibility that exercise induced left axis shifts may be associated with obstructive disease of the left anterior descending artery. First the blood supply to the left anterior fasciculus as described by Rosenbaum and associates¹⁷ is derived entirely from the left anterior descending artery. However it is not clear if this arrangement persists in a diseased heart with multiple collateral vessels.¹⁸ Second in Rosenbaum's series the most common cause of left anterior fasciculus block was

thought to be coronary artery disease. And third transient left anterior fasciculus block is sometimes found in association with an acute anteroapical myocardial infarction.

Of the 4 patients in the Minnesota series who developed exercise induced left axis shifts 3 developed a shift beyond -45 degrees consistent with a complete left anterior fasciculus block and the fourth developed a shift to -16 degrees consistent with a partial left anterior fasciculus block. The finding that 3 of the 4 patients in this group did have complete obstruction of

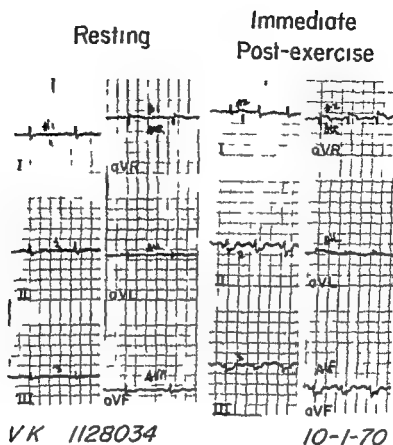


Fig 6 Pre and postexercise ECGs of V K (extremity leads only) showing a transient exercise-induced left axis shift prior to coronary artery surgery

the left anterior descending artery is very suggestive of a causal relationship. However, the fourth patient in this group had normal coronary arteriograms. And since there were only 4 patients in the entire group, it was not possible to make a satisfactory comparison to the total group of patients analyzed. Hence the significance of the apparent association between exercise induced left axis shifts and severe obstruction of the left anterior descending artery remains unclear.

It also seems possible that exercise induced right axis shifts could be associated with obstructive disease of the right and left coronary arteries. However on the basis of Rosenbaum's work it would seem that such an association would be found less frequently with right axis shifts than with left axis shifts. First, the blood supply to the left posterior fasciculus is derived from both the left and the right coronary arteries. Second, the left posterior fasciculus is both shorter and thicker than the left anterior fasciculus. Third, left posterior fasciculus blocks occur with a much lower

Table VI Frequency of at least 50 per cent obstruction in patients with exercise induced axis shift

| | No of patients | Less than 50% obstruction | At least 50% obstruction |
|-----------------------|----------------|---------------------------|--------------------------|
| Left axis shift | 4 | 1 | 3 |
| Right axis shift | 9 | 5 | 4 |
| All patients analyzed | 154 | 51 | 103 |

frequency than left anterior fasciculus block.

Of the 9 patients in this series who developed exercise induced right axis shifts only one developed a shift beyond +120 degrees and only 3 developed shifts beyond +100 degrees. Thus the postexercise axis approached Rosenbaum's criteria of about +120 degrees for complete left posterior fasciculus block in only 3 of the 9 patients. There was no consistent association between the localization of severity

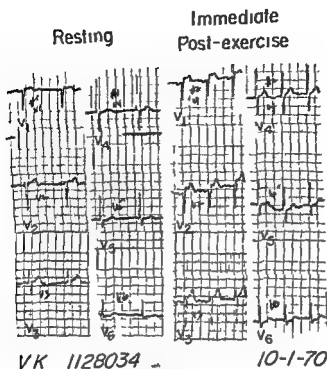


Fig 7 Pre and postexercise ECG's of V K (precordial leads only) showing transient S-T-segment elevations in Leads V_1 to V_5 prior to coronary artery surgery

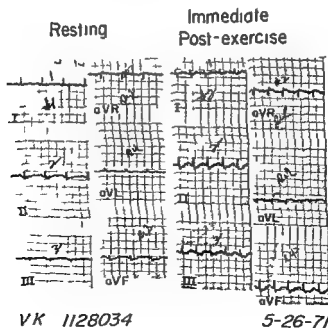


Fig 8 Pre- and postexercise ECG's of V K (extremity leads only) showing a reversal of the preoperative exercise-induced left axis shift after a successful surgical bypass of an obstructing lesion in the left anterior descending artery

obstruction was greater than 85 per cent in 16 patients and greater than 50 per cent in a seventeenth patient. The remaining patient had a normal coronary arteriogram and the most minimal exercise induced ST-segment elevations. Nine of the 158 patients had exercise induced right axis shifts. Only 4 of these 9 patients had greater than 50 per cent obstruction of a major coronary artery as compared to 103 of 154 patients in the total group studied. Also there was no trend toward predominant involvement of any particular coronary artery in these 4 patients. Hence it appears that this finding is not predictably associated with severe localized coronary artery disease. Only 4 of the 158 patients had exercise induced left axis shifts. Three of these four patients had complete obstruction of the left anterior descending artery. But the group size was small and the fourth patient had a normal coronary arteriogram. Hence it is only possible to suggest that this finding may be associated with severe disease of the left anterior descending artery. The results of coronary artery surgery are discussed in a patient who had both a left axis shift and precordial ST-segment elevations on his preoperative exercise ECG.

REFERENCES

- 1 Masters A M and Rosenfeld I Two-step exercise test. Current status after twenty five years. *Mod Concepts Cardiovasc Dis* 36:19 1967
- 2 Robb G M and Marks H H Post-exercise electrocardiogram in arteriosclerotic heart disease. *JAMA* 200:918 1965
- 3 Doyle J T and Kinch M S The prognosis of an abnormal electrocardiographic stress test. *Circulation* 41:545 1970
- 4 Burchell H B The value of exercise tests in the diagnosis of coronary disease in Symposium on Coronary Heart Disease ed 2 Monograph No 2 1968 American Heart Association p 49
- 5 Prinzmetal M Kenamer R Merliss R et al Angina pectoris I A variant form of angina pectoris. *Am J Med* 27:375 1959
- 6 Silverman M E and Flamm M D Variant angina pectoris. *Ann Intern Med* 75:339 1971
- 7 Fortuin N J and Friesinger C C Exercise-induced ST segment elevation. *Am J Med* 49:159 1970
- 8 Doan A E Peterson D R Blackmon J R and Bruce R A Myocardial ischemia after maximal exercise in healthy men. A method for detecting potential coronary heart disease. *Am Heart J* 69:111 1965
- 9 Bayley R H and LaDue J S Electrocardiographic changes of impending infarction and the ischemia injury pattern produced in the dog by total and subtotal occlusion of a coronary artery. *Am Heart J* 28:154 1944
- 10 Wegeris R Segers M heating R P and Ward H P Relationship between the reduction in coronary flow and the appearance of electrocardiographic changes. *Am Heart J* 38:90 1949
- 11 Gorlin R Klein M D and Sullivan J M Prospective correlative study of ventricular aneurysm. *Am J Med* 42:519 1967
- 12 Rubenstein C Yattreau R and Walston A Significance of exercise ST elevation. *Circulation* 43:44 (Suppl 11) 219 1971
- 13 Fernandez R Scobal L and Lenegre J Electrocardiography study of left intraventricular hemiblock in man during selective coronary arteriography. *Am J Cardiol* 26:1 1970
- 14 Wagner R and Rosenbaum M Transient left posterior hemiblock Association with acute lateral myocardial infarction. *Am J Cardiol* 29:558 1972
- 15 Bruce R A Detry J M Early K and Early R Polarcardiographic responses to maximal exercise in healthy young adults. *Am Heart J* 70:206 1972
- 16 Kulbertus H E and Humblet L Transient hemiblock. An abnormal type of response to the Master two-step test. *Am Heart J* 83:574 1972
- 17 Rosenbaum M B Elizari M V and Lazzari J O The hemiblocks. New concepts of intraventricular conduction based on human anatomical physiological and clinical studies. *Oldsmar Fla* 1970 Tampa Tracings
- 18 James T N Anatomy of the coronary arteries. New York 1961 Paul H Hoeber Inc Medical Division of Harper & Row Publishers

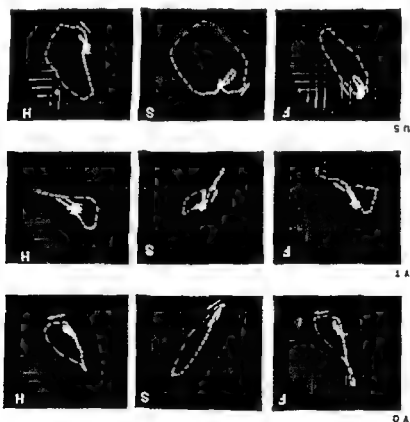


Fig 2 Vectorcardiograms of three patients demonstrating intraventricular conduction defects of varying degrees. F = frontal plane. S = sagittal plane. H = horizontal plane.

Table 1 Electrophysiological findings

| Patient | Age yr | Sex | ECG | | | | | VCG |
|---------|--------|-----|--------------------|--------------------|-----------------|-------|--------|------------------|
| | | | P-R interval (sec) | QRS duration (sec) | Electrical axis | Low P | Others | |
| | | | 4 H (misc) | H V (misc) | | | | Frontal QRS loop |

| | | | | | | | | | |
|-----|----|---|------|------|------|-------------------|-----|----|---|
| Y O | 37 | M | 0.32 | 0.14 | -50° | + Bradycardia | 128 | 88 | Early forces directed left in frontal plane with main forces superiorly |
| Y T | 34 | M | 0.24 | 0.09 | +80° | + Incomplete RBBB | 160 | 44 | Terminal delay |
| U S | 45 | M | 0.20 | 0.08 | +75° | - | 100 | 56 | Slight terminal delay |

the same right femoral vein where a Damato hexapolar probe had been previously introduced. The cutting claw was advanced to the apex of the right ventricle under fluoroscopic observation. One of the biopsy specimens was fixed in ten per cent formaldehyde solution and stained with the hematoxylin and eosin. For electron micro-

scopic examinations an additional specimen was immediately fixed with two per cent phosphate buffered osmium tetroxide for two hours. After dehydration with increasing concentrations of alcohol the tissue was embedded in araldite VI and sectioned with a microtome. Thin sections were stained with uranylacetate alone and

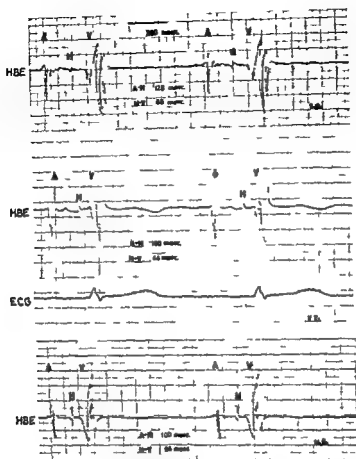


Fig 3 His bundle electrograms from three patients showing atrial (A) His bundle (H) and ventricular (V) potentials and AH and HV intervals. Simultaneous recording of HBE and ECG is shown in the middle (patient Y T)

in combination with lead citrate. Micrographs were taken with a JEM 7 electron microscope.

Results

Physical examination of the heart revealed no change among the three patients. None of the three claimed to have experienced symptoms of angina pectoris. X-ray examination of the chest revealed minimum cardiomegaly in patient Y O. Moderate hypotension was present in all three cases. Two cases (patient Y O and patient Y T) were found to have electrocardiographic abnormalities as shown in Fig 1 and as listed in Table I.

Vectorcardiographic abnormalities were found definitely in two cases (Fig 2). In patient Y O the frontal QRS loop

presented initial forces directed left inferiorly with main forces directed superiorly and the delay of the terminal portion. The patient's ECG showed left axis deviation (-50 degrees), the prolonged PQ interval (0.32 sec), the prolonged QRS complex (0.14 sec) and the deep S in V_1 and V_6 . Conduction delay of the terminal portion of the loop was also found right posteriorly in patient Y T whose standard ECG showed rSR in V_1 and a slightly prolonged QRS complex. In patient U S the vectorcardiographic tracings revealed slight conduction delay of the terminal portion.

Tracing of the HBE was made at one slow sinus rate. Electrical activity of the His bundle appeared as a biphasic or triphasic wave occurring between the atrial and ventricular electrograms. Fig 3 illus-

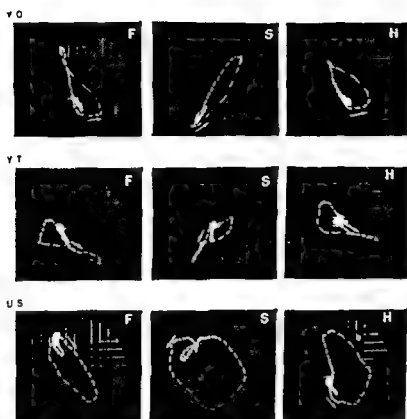


Fig 2 Vectorcardiograms of three patients demonstrating intraventricular conduction defects of varying degrees F = frontal plane S = sagittal plane H = horizontal plane

Table I Electrophysiological findings

| Patient | Age yr | Sex | ECG | | | | | HBE | | VCG |
|---------|-----------|-----|-------------------------|--------------------------|--------------------|----------|-----------------|--------------------------|--------------------------|---|
| | | | PR interval (sec) | QRS duration (sec) | Electrical axis | Low P | Others | AH interval (msec) | HV interval (msec) | Frontal QRS loop |
| YO | 37 | M | 0.32 | 0.14 | -50 | + | Bradycardia | 128 | 88 | Early forces directed left inferiorly with main force superiorly terminal delay |
| YT | 34 | M | 0.24 | 0.09 | +80 | + | Incomplete RBBB | 160 | 44 | Terminal delay |
| US | 45 | M | 0.20 | 0.08 | +75 | - | | 100 | 56 | Slight terminal delay |

the same right femoral vein where a Damato hexapolar probe had been previously introduced. The cutting claw was advanced to the apex of the right ventricle under fluoroscopic observation. One of the biopsy specimens was fixed in ten per cent formaldehyde solution and stained with the hematoxylin and eosin. For electron micro-

scopic examinations an additional specimen was immediately fixed with two per cent phosphate buffered osmium tetroxide for two hours. After dehydration with increasing concentrations of alcohol the tissue was embedded in araldite and sectioned with a microtome. Thin sections were stained with uranylacetate alone and

V 1 mR 86
V mR 5

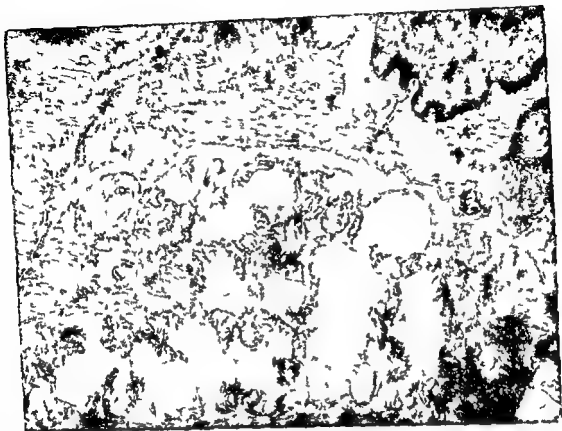


Fig 5 Electron micrograph (patient Y O) Numerous degenerated mitochondria of which membrane is double on one side but single on the opposite side. Granular substance appears to have bulged out into the outer space and to be surrounded by the inner and outer membrane (Original magnification $\times 10,000$)

trates the HBE obtained from the three patients. Simultaneous recordings of the HBE and Lead II ECG are shown in the middle (patient Y T). The A-H interval of the three patients were 128, 160 and 100 msec and the H-V interval 88, 44 and 56 msec respectively.

The light microscopic photographs of biopsied myocardium (Fig 4) showed wide variations of heart muscle cells in size and massive infiltration of fatty tissue. The nuclei were uneven and some of them were rectangular. Alterations in the arrangement of muscle fibers were evident on the longitudinal sections. Focal vacuolar degeneration of myocardial cells was seen in transverse sections. There was neither fibrosis nor increase in cellularity apart from scattered focal hemorrhages. Endomyocardial fibrosis and fibroelastosis were not found.

The prominent myocardial changes revealed by electron microscopy in patient

Y O were the degeneration of the mitochondria with disorganization of internal structure. Degenerated mitochondria were bounded by double limiting membrane on one side but by single membrane on the opposite side. Granular substance which was less electron dense was found scattered in the outer space (Fig 5). The majority of mitochondria, however, appeared to be essentially normal in patient U S. Myofibrillar alterations characterized by disarray and dissolution of the Z line and I band were often found. There was also focal contractile element damage with disorganization and loss of myofilaments. The external lamina generally appeared to be intact.

Patient Y O who had been treated for neurological rehabilitation in Kagoshima University Kirishima Branch Hospital was taken to our department because of syncopal episodes with bradyarrhythmia.



Fig 4 A and B A Light micrograph of myocardium of right ventricle of patient U S Note wide variation of muscle fibers in size and massive involvement of fatty tissue The nuclei are uneven and some of them are rectangular (Hematoxylin and eosin Original magnification A $\times 160$ B $\times 400$)

conduction system in a case showing atrial flutter with varying degrees of A V block but found no anatomic basis for the electrocardiographic abnormalities. In Thomson's study¹¹ the S A node was involved by virtue of the site of the atria and the A V node was also affected though to a lesser extent than the S A node. Furthermore it is worthy to mention that the bundle branch was involved in an extensive area of myocardial change. This denotes the prolongation of H V interval which means the conduction disturbances distal to the His bundle. Intraventricular conduction defects of varying degrees were also shown by vectorcardiographic tracings. Conduction delay of the terminal portions was oriented superiorly from the point of origin.

In 1964 Petkovich and colleagues¹⁰ reported a case of myotonic dystrophy that developed Stokes-Adams syndrome due to A V dissociation. An implantation of a pacemaker allowed them to carry out myocardial biopsy taken from the pericardium and they found rather indistinctly bordered myocardial muscle fibers feathery cytoplasm and longitudinal separation of myofibrils. In postmortem cases microscopic changes of the heart were fatty infiltration of the myocardium, interstitial fibrosis of varying cellularity and sometimes with lymphocytic infiltration.^{13, 14, 15} The myocardial findings of our cases were all most similar to theirs except for cellular involvement. It has been accepted that the classical histological finding of the hearts in myotonic dystrophy was non-specific.^{17, 18}

Recently Bulloch and co-workers²² reported electron microscopic changes of the left ventricular muscles of a patient with marked cardiomegaly. The most frequent myocardial abnormalities were found in the sarcoplasmic reticulum and mitochondria in cells that otherwise appeared intact. In our cases the most striking abnormalities were found in the mitochondria. The internal structure was lost and a peculiar substance which was granular appeared to have bulged into the outer space. It was not always possible to be sure whether this finding might be a primary site of pathologic change. Serial biopsy studies in various stages of the disease are necessary for the detection of early abnormality.

Örndahl and colleagues⁴ found out by vector and scalar-cardiographic studies that the diffuse change of the specialized conduction system was more dominant than the disorganization of the contractile elements of the myocardium. Our results of examination using light and electron microscopy of the biopsied myocardium showed the presence of distinctive involvement of the myocardium. It seems therefore likely that both the conduction system and the myocardium are extensively involved in an advanced stage of myotonic dystrophy.

The pacemaker implanted in patient Y O has been working well for more than twelve months, but we are pessimistic about this patient's future because of the involved serious cardiomyopathy as revealed by biopsy. His younger sister who was suspected to have myotonic dystrophy died accidentally when she was asleep five days after an operation for cataract. We had still another patient who died suddenly while waiting for hospitalization. Other cases of an unexpected death^{11, 12} were also reported. Careful attention should be paid not only to neuromuscular defects and infections but also to cardiac lesions in myotonic dystrophy.

Summary

For frequent complications of cardiac involvement it is important to recognize exactly myotonic dystrophy from a cardiovascular standpoint. Electrophysiological and histological studies were performed in three patients. Analysis of the site of the conduction disturbances was revealed by His bundle electrography proximal to the His bundle in one patient and distal in two other patients. The histological examination for biopsied myocardium by light and electron microscopy disclosed replacement of large areas by fat and also showed varying degrees of cellular damage. The most striking abnormality of the heart muscles was found in the mitochondria in which a peculiar and granular substance bulged into the outer space. Both the conduction system and the myocardium were involved in our cases. The value of His bundle recordings and endomyocardial biopsy in evaluating cardiac abnormality in the disease were discussed.

Eight days later he developed Stokes Adams syndrome due to cardiac arrest for more than three minutes, from which he recovered with the administration of intra venous isoproterenol. On his twenty third hospital day, the electrode of a permanent cardiac pacemaker (ventricular inhibited type) was anchored to the epicardium over the outflow tract of the right ventricle. Seven days postoperatively, it was found that a segment of the skin overlying the pace generator was necrotic. An ECG made after the implantation of the pacemaker showed natural, fusion, and pacemaker induced beats. The induced QRS complexes had the appearance of those of left bundle branch block.

Discussion

Myotonic dystrophy, originally described by Steinert,¹⁰ is a heredo familiar neuro muscular disease in which cardiac involvement is common and serious. In most cases, neuromuscular defects are apparent long before any evidence of the heart disease but in a few cases the latter may be the predominant feature and an important aid to the early diagnosis.^{2,11} Griffith¹ provided a clinical description of a patient with a heart rate of 46 to 30 per minute and was the first to call attention to the disturbances of cardiac rhythm in association with this disease. Since then some investigators have described cardiac manifestations in myotonic dystrophy. There were few clinical signs of the heart disease the most prominent findings being electro cardiographic abnormalities such as various kinds of arrhythmias and conduction defects.^{4,12} The incidence of electrocardiographic abnormalities was reportedly found in 40 to 85 per cent, but cases of complete atrioventricular dissociation with Stokes Adams attacks were rarely documented.^{13,14} The pathogenesis of cardiac alterations, however, remains obscure. To our knowledge the present communication is the first in reporting the use of the His bundle electrogram for the determination of the site of conduction disturbances and endomyocardial biopsy for the histological investigations of the abnormalities of contractile elements.

The PR interval was found prolonged for more than 0.2 sec in two cases. Payne

and Greenfield¹⁵ reported prolongation of the PR interval in five of 21 cases and Örndahl and associates⁴ in 11 of 29 patients. The standard electrocardiogram however, fails to localize the site of delay or block precisely because the PR interval merely represents the sum of intra atrial conduction time through the A V node the His bundle, the bundle branches and a part of the Purkinje fibers. The technique of HBE recently developed by Damato and colleagues¹⁶ have contributed to our understanding of the modes of transmission of the impulses from the atrium to the ventricle. Based upon the normal range reported by Narula and associates¹⁷ prolongation of the A H interval was found in patients Y T and Y O, and that of the H V interval in patients Y O and U S. According to Rosen and co workers,¹⁸ most patients with the prolongation of the PR interval and the intraventricular conduction defects had the prolongation of P H (the interval from the P wave to the first high frequency component of the H wave) suggesting the lesion proximal to the His bundle. In patient Y O whose QRS duration was also prolonged both A H and H V intervals were prolonged. The latter was more prolonged and the major site of delay was suspected to be distal to the His bundle. In fact he soon developed Stokes Adams attacks due to simultaneous atrial and ventricular standstill. The lesion was suspected to be proximal to the His bundle in patient Y T. In patient U S, the H V interval was found prolonged although his PR interval and QRS complex were normal. This finding suggests that there was a first degree block in the distal portion of the main stem of the His bundle below the site of His bundle recording. It seems likely that not only the proximal but also the distal region to the His bundle was involved in myotonic dystrophy. When first seen, all cases were found progressed showing typical features of the disease so the early cases may allow us to determine whether progression of conduction disturbances occurs either proximally or distally to the His bundle.

Histological changes in the specialized conduction system in myotonic dystrophy were reported by a few investigators. Cannon¹⁷ attempted to study the lesion of the

Immunologic findings in idiopathic cardiomyopathy

A prospective serial study

Allan B Karsner MD*

Exelyn V Hess MD FACP**

Noble O Fowler MD FACP***

Cincinnati Ohio

Idiopathic cardiomyopathy (IC) is a disease of unknown etiology with suspected immunologic mechanisms.^{1,2} Serum factors reactive to heart muscle constituents have been reported in between 11 and 30 per cent of sera from patients with IC.³⁻⁶ Bound gamma globulin^{7,8} and complement (C3)⁹ have been found in the ventricular muscle of patients who died with this disease the hearts of control subjects being free of such deposits. Previous studies have not clarified whether these findings were primary and related to etiology and/or pathogenesis or secondary and represented a nonspecific immunologic response to heart muscle damage.

In many diseases with immunologic mechanisms serologic factor titers vary and are often correlated with disease activity. In systemic lupus erythematosus where renal and skin manifestations are related to the deposition of antibody antigen complexes the titer of anti DNA antibody parallels disease activity and during a remission the antibody response may be

absent or in a low titer.¹⁰ Serum complement levels fall just prior to and in relation to an exacerbation of renal disease in systemic lupus erythematosus.¹¹

Serial studies of sera in acute rheumatic fever have shown heart reactive factors (H R F) in 25 to 63.4 per cent of patients and their presence has been correlated with the development and recurrence of clinical symptoms.¹²⁻¹⁴ Bound gamma globulin has been found in 18 per cent of auricular biopsy specimens from patients with clinically inactive rheumatic heart disease.¹⁵ Sixteen per cent of patients with chronic rheumatic heart disease were found to have H R F.¹⁶

We have previously reported the results of an immunologic study of patients with IC.¹ Thirty three patients were subjected to a single study but the data did not show a high incidence of serologic immune factors. The present study was undertaken to evaluate serially the presence of humoral immune factors in a group of patients with idiopathic cardiomyopathy and to attempt

*From the Divisions of Cardiology and Immunology Department of Internal Medicine University of Cincinnati College of Medicine Cincinnati Ohio

Supported in part by National Institutes of Health Grant HL 4307 HL 5445 of the United States Public Health Service and Research Grant HL 4307 HL 5445 of the United States Public Health Service

Received for publication June 9, 1973

Reprint requests to Allan B Karsner MD 4235 Soro Rd Toledo Ohio 43623

**Clinical Assistant Professor of Medicine Medical College of Ohio Toledo

***Professor of Medicine Director Division of Immunology University of Cincinnati College of Medicine

The authors wish to thank Pro Dr A Igata for his encouragement and also extend their thanks to Dr K Kawamura the Third Department of Internal Medicine Faculty of Medicine Kyoto University for his valuable advice in the electron microscopic studies. The implantation of the permanent pacemaker was done by Dr A Taira the Second Department of Surgery Faculty of Medicine Kagoshima University.

REFERENCES

- 1 Griffith T W On myotonia Q J Med 5:229 1911
- 2 Evans W The heart in myotonia atrophica Br Heart J 6:41 1944
- 3 DeWind L T and Jones R J Cardiovascular observations in dystrophic myotonic JAMA 144:299 1950
- 4 Örndahl G Thuleus O Enestrom S and Delhin O The heart in myotonic disease Acta Med Scand 176:479 1964
- 5 Spillane J D The heart in myotonia atrophica Br Heart J 13:343 1951
- 6 Waring J Ravin A and Walker C Studies in dystrophia myotonica II Clinical features Arch Intern Med 65:763 1940
- 7 Frank E An accurate clinically practical system for spatial vectorcardiography Circulation 19:737 1956
- 8 Scherlag H J Lau S H Helfant R H Berkowitz W D Stein E and Damato A N Catheter technique for recording His bundle activity in man Circulation 39:13 1969
- 9 Konno S and Sakakibara S Endomyocardial biopsy Dis Chest 44:345 1963
- 10 Steinert H Über das klinische und anatomische Bild des Muskelschwundes der Myotoniker Dtsch Z Nervenheilkd 37:58 1909
- 11 Holt J M and Lambert E H N Heart disease and the presenting feature in myotonia atrophica Br Heart J 26:433 1964
- 12 Spurny O and Wolf J Prolonged atrial flutter in myotonic dystrophy Am J Cardiol 10:886 1962
- 13 Payne C A and Greenfield J C Electrocardiographic abnormalities associated with myotonic dystrophy AM HEART J 6:436 1963
- 14 Damato A N Lau E H Helfant R H Stein E Berkowitz W D and Cohen S H Study of atrioventricular conduction in man using electrode catheter recordings of His bundle activity Circulation 39:287 1969
- 15 Narula O S Scherlag B J Samet P and Javier R P Atrioventricular block Localization and classification by His bundle recordings Am J Med 50:146 1971
- 16 Rosen K M Rahimtoola S H Chuquimia R Loeb H S and Gunnar R M Electrophysiological significance of first degree atrioventricular conduction disturbance Circulation 43:491 1971
- 17 Cannon P J The heart and lungs in myotonic muscular dystrophy Am J Med 32:765 1962
- 18 Thomson A M P Dystrophia cordis myotonica studied by serial histology of the pacemaker and conducting system J Pathol Bacteriol 96:285 1968
- 19 Petkovich N J Dunn M and Reed W Myotonia dystrophica with AV dissociation and Stokes Adams attacks A case report and review of literature AM HEART J 68:391 1964
- 20 Litchfield J A AV dissociation in dystrophia myotonica Br Heart J 15:357 1953
- 21 Fisch C and Evans P V The heart in dystrophia myotonica report of an autopsied case N Engl J Med 251:527 1954
- 22 Bulloch R T Davis J L and Hara M Dystrophia myotonica with heart block A light and electron microscopic study Arch Pathol 84:130 1967

Table I Immunologic data and temporal correlation of positive tests to congestive heart failure in 32 patients with idiopathic cardiomyopathy

| Test | Positive | Negative | Temporal correlation of positive tests to cardiac decompensation |
|-------------------------------|----------|----------|--|
| H.R.F. | 9 | 23 | 2 |
| ANA | 0 | 32 | 0 |
| Latex test for RF | 9 | 23 | 2 |
| SSCAT | 0 | 32 | 0 |
| Depression of complement (C3) | 1 | 31 | 1 |
| Elevation of ASO titer | 3 | 29 | 0 |
| ATA | 6 | 26 | 2 |

Abbreviations: H.R.F. = heart failure; A.A. = anti-leaf (C3); R.F. = rheumatoid factor; SSCAT = sensitized sheep cell agglutination test; ASO = antistreptolysin O; ATA = thyroid antiodies

Other serologic tests The following procedures were performed on all sera: (1) rheumatoid factor (anti-gammaglobulins) by the Latex Fixation Test (Hyland Slide Test) and Sensitized Sheep Cell Agglutination Test (SSCAT) (2) standard tests for syphilis (3) total serum proteins with albumin globulin ratio (4) serum immunoelectrophoresis (I.E.P.A.) (5) thyroid antibodies by the thyroglobulin agglutination test (Hyland Slide Test) and indirect immunofluorescent test for microsomal antibodies (6) anti-nuclear factor by an indirect immunofluorescent test on infant thyroid (7) Antistreptolysin O (ASO) titer and (8) serum enzyme battery including lactic dehydrogenase, creatinine phosphokinase and glutamic transaminase.

The Latex Slide Test was read on a scale of 1+ to 4+ and only tests that are 2+ or greater are reported as positive. The I.E.P.A. was read on a scale of 1+ to 4+ by a single individual. Only elevations or depressions of 2+ or greater are reported as positive.

All serologic tests were performed 3 to 4 times a year during periods of stable heart disease at the onset of exacerbation of congestive heart failure and at more frequent interval during severe congestive heart failure.

Results

Nine of the 32 patients had positive H.R.F. tests and in eight of the nine it was present on more than one occasion. Two patients demonstrated sarcoid-like subarcoid (SS) staining; one patient showed

only intermyofibrillar staining (I.I.F.) and six patients showed both SS and I.I.F. staining. In two of the nine patients there was a correlation between the appearance of H.R.F. and exacerbation of the patient's congestive heart failure. In the other seven patients there was no correlation. Three patients demonstrated elevated ASO titers but in none of these was H.R.F. present. Hyperglobulinemia was noted in eight of the 32 patients but in only one patient did the appearance of hyperglobulinemia correlate with an exacerbation of the patient's congestive heart failure. Anti-nuclear antibodies were not detected. Nine of the 32 patients had positive latex slide tests during the course of the disease and in two of these the appearance of the positive latex test correlated with an exacerbation of the patient's congestive heart failure. No patient with a positive latex test had a positive SSCAT. Six of the 32 patients had positive antithyroid antibody tests and in two of these there was correlation with the onset of congestive heart failure. Two had a positive standard test for syphilis (V.D.R.L.), one being a biologic false positive. Elevated serum enzymes were noted in nine of 32 patients and in two of these there was correlation with increasing congestive heart failure. One patient had a preterminal depression of C3. These results are summarized in Table I.

Table II correlates the positive serologic findings with changes in the patient's clinical status. Five of the 18 patients with stable heart disease had positive H.R.F.

to correlate changes in immunologic variables with exacerbations and remissions of disease. The serologic factors studied included anti-gammaglobulin and anti-nuclear factors (ANA), thyroid antibody (ATA), immunoglobulin and complement levels, and serum HRF.

Materials and methods

Patient selection The diagnosis of idiopathic cardiomyopathy in many respects is a diagnosis of exclusion. Patients entered into this study were evaluated 3 to 4 times a year with a physical examination, chest x-ray, and electrocardiogram. Periodic phonocardiograms, vectorcardiograms, and cardiac fluoroscopies were performed, and several of the patients had cardiac catheterization studies. All patients demonstrated cardiac enlargement at some time during their clinical course. Each patient had either a ventricular or atrial gallop or a summation gallop during periods of congestive heart failure. All patients had abnormal electrocardiograms. No patient was included who had a history of angina pectoris, myocardial infarction, or electrocardiographic evidence of previous myocardial infarction. Patients with a sustained blood pressure of 150/100 mm Hg or more were considered to have hypertensive cardiovascular disease and were not included in this study. A diastolic cardiac murmur, valvular calcification, diabetes mellitus, hypercholesterolemia, or hyperthyroidism excluded patients from this study. No patient had evidence of idiopathic hypertrophic subaortic stenosis. A total of 38 patients satisfied these criteria and were placed in the prospective study. Three patients were lost to follow-up and three subsequently showed evidence of another etiology for their heart disease.

Thirty-two patients were followed for a period of 51.6 patient-years. Twenty-two of these were observed for over 1 year and 11 were observed for over 2 years. Of the 32 patients, four died and the three who were autopsied had findings consistent with idiopathic cardiomyopathy. Ten had an exacerbation of disease with new or more severe congestive heart failure on physical examination and cardiac enlargement on chest x-ray. Eighteen had stable heart dis-

ease. There were 20 men and ten women. Twenty-four patients were black and eight were white. The mean age at onset was 36.5 years. All patients were questioned as to possible triggering mechanism of the first bout of congestive heart failure. Seven had a daily alcohol intake that exceeded two ounces of hard liquor per day. Six had a flu-like syndrome prior to the onset of congestive heart failure. Five developed congestive heart failure during the first three months postpartum. One developed congestive heart failure following a flu-like syndrome and also had a history of alcoholic intake. No triggering mechanism was elicited from the remaining 12 patients.

Heart tissue Samples of right ventricular muscle were obtained during cardiac surgery from a patient with tetralogy of Fallot. These were quickly frozen in liquid nitrogen and stored at -40°C . The tissue was tested for antigenicity with known positive and negative sera before being used in this study.

Antisera Goat antisera to human immunoglobulins was prepared and conjugated with fluorescein isothiocyanate by the methods previously described.¹⁰

Staining procedure Tissue sections were cut at 5 micra, fixed in acetone for 10 minutes at room temperature, air dried, and rinsed in buffered saline and incubated with test serum for 1 hour. The slides were then washed in buffered saline, incubated with conjugated goat anti-human gamma globulin for 1 hour, again washed in buffered saline, and mounted in buffered glycerol. Each run contained positive and negative control sera. Less than 5 per cent of normal sera give positive reactions. The sections were examined with an American Optical Fluorostar Microscope equipped with an Osram HBO 200 Mercury Lamp, 2 mm Corning 5840 filter, and a red excluding BG 14 filter. The slides were read independently by two observers without knowledge of the source of the serum being tested. The degree of fluorescence was graded on a 0 to 4+ scale. 2+ fluorescence being required for a positive reading. All positive sera were rerun in dilution. Three patterns of immunofluorescent staining of cardiac myofibers were seen: sarcolemmal subsarcolemmal (SS), intermyofibrillar (IMF), and diffuse (D).

Table I Immunologic data and temporal correlation of positive tests to congestive heart failure in 32 patients with idiopathic cardiomyopathy

| Test | Positive | Negative | Temporal correlation of positive tests to cardiac decompensation |
|-------------------------------|----------|----------|--|
| HRF | 9 | 23 | 2 |
| ANA | 0 | 32 | 0 |
| Latex test for RF | 9 | 23 | 2 |
| SSCAT | 0 | 32 | 0 |
| Depression of complement (C3) | 1 | 31 | 1 |
| Elevation of ASO titer | 3 | 29 | 0 |
| ATA | 6 | 26 | 2 |

Abbreviations: HRF = heart factor test; ANA = antinuclear factor; RF = rheumatoid factor; SSCAT = sensitized sheep cell agglutination test; ASO = antistreptolysin O; ATA = thyroid antithyroid antibody.

Other serologic tests The following procedures were performed on all sera: (1) rheumatoid factor (anti-gammaglobulins) by the Latex Fixation Test (Hyland Slide Test) and Sensitized Sheep Cell Agglutination Test (SSCAT)¹⁴; (2) standard tests for syphilis; (3) total serum proteins with albumin globulin ratio; (4) serum immunoelectrophoresis (IEPA); (5) thyroid antibodies by the thyroglobulin agglutination test (Hyland Slide Test) and indirect immunofluorescent test for microsomal antibodies¹⁵; (6) anti-nuclear factor by an indirect immunofluorescent test on infant thyroid; (7) Antistreptolysin O (ASO) titer and (8) serum enzyme battery including lactic dehydrogenase, creatinine phosphokinase and glutamic transaminase.

The Latex Slide Test was read on a scale of 1+ to 4+ and only tests that are 2+ or greater are reported as positive. The IEPA was read on a scale of 1+ to 4+ by a single individual. Only elevations or depressions of 2+ or greater are reported as positive.

All serologic tests were performed 3 to 4 times a year during periods of stable heart disease at the onset of exacerbation of congestive heart failure and at more frequent intervals during severe congestive heart failure.

Results

Nine of the 32 patients had positive HRF tests and in eight of the nine it was present on more than one occasion. Two patients demonstrated sarcolemmal subarcuolar (SS) staining; one patient showed

only intermyofibrillar staining (IMF) and six patients showed both SS and IMF staining. In two of the nine patients there was a correlation between the appearance of HRF and exacerbation of the patient's congestive heart failure. In the other seven patients there was no correlation. Three patients demonstrated elevated ASO titers but in none of these was HRF present. Hyperglobulinemia was noted in eight of the 32 patients but in only one patient did the appearance of hyperglobulinemia correlate with an exacerbation of the patient's congestive heart failure. Anti-nuclear antibodies were not detected. Nine of the 32 patients had positive latex slide tests during the course of the disease and in two of these the appearance of the positive latex test correlated with an exacerbation of the patient's congestive heart failure. No patient with a positive latex test had a positive SSCAT. Six of the 32 patients had positive antithyroid antibody tests and in two of these there was correlation with the onset of congestive heart failure. Two had a positive standard test for syphilis (VDRL), one being a biologic false positive. Elevated serum enzymes were noted in nine of 32 patients and in two of these there was correlation with increasing congestive heart failure. One patient had a preterminal depression of C3. These results are summarized in Table I.

Table II correlates the positive serologic findings with changes in the patient's clinical status. Five of the 18 patients with stable heart disease had positive HRF

Table II Correlation of positive serologic tests with changes in the patient's clinical status in 32 patients with idiopathic cardiomyopathy

| Test | 18 patients with stable heart disease | 10 patients who developed CHF* | 4 patients who died |
|-------------------------------|---------------------------------------|--------------------------------|---------------------|
| HRF | 5 | 3 | 1 |
| ANA | 0 | 0 | 0 |
| Latex test for RF | 4 | 3 | 2 |
| SSCAT | 0 | 0 | 0 |
| Depression of complement (C3) | 0 | 0 | 1 |
| ATA | 2 | 3 | 1 |

*CHF = congestive heart failure. Other abbreviations same as in Table I.

Table III Heart reactive factor titers in 9 patients with idiopathic cardiomyopathy

| Patient No | Cardiac status | Titer of heart reactive factor | | | |
|------------|-----------------|--------------------------------|-----|------|------|
| | | 1:1 | 1:5 | 1:10 | 1:20 |
| 1 | Borderline CHF* | 1+ | 1+ | 0 | 0 |
| | Severe CHF | 2+ | 1+ | + | 0 |
| 2 | Borderline CHF | 2+ | 1+ | + | 0 |
| | Mild CHF | 2+ | 1+ | 0 | 0 |
| 3 | Compensated | 2+ | 1+ | + | 0 |
| | Moderate CHF | 2+ | + | 0 | 0 |
| 4 | Compensated | + | 0 | 0 | 0 |
| | Mild CHF | 3+ | 1+ | 0 | 0 |
| 5 | Borderline CHF | 2+ | 0 | 0 | 0 |
| | Borderline CHF | 2+ | 0 | 0 | 0 |
| 6 | Compensated | 2+ | 1+ | 0 | 0 |
| | Compensated | 2+ | 0 | 0 | 0 |
| 7 | Acute onset CHF | 0 | 0 | 0 | 0 |
| | Compensated | 2+ | 0 | 0 | 0 |
| 8 | Compensated | 1+ | 0 | 0 | 0 |
| | Compensated | 2+ | 0 | 0 | 0 |
| 9 | Compensated | 2+ | 0 | 0 | 0 |
| | Compensated | 2+ | 0 | 0 | 0 |

*CHF = congestive heart failure.

four had positive latex tests and two had positive thyroid antibody tests. Three of ten patients with increasing congestive heart failure had positive HRF, and in two there was correlation between the appearance of HRF and an exacerbation of heart failure. Three had positive latex tests and three had thyroid antibodies; the appearance of these positive tests usually did not correlate with a change in the patient's clinical condition. One of the

four patients who died had a positive HRF but there was no correlation between its appearance and exacerbation of the patient's disease.

Table III demonstrates the HRF titers on the nine positive patients and correlates these with the patient's clinical condition. HRF is present in low titer in patients with IC and a meaningful correlation does not appear to exist between titer of HRF and severity of the disease.

There was no correlation between positive latex tests and positive tests for HRF or between positive thyroid antibody and HRF tests.

Of the nine patients with HRF four had a daily alcohol intake that exceeded two ounces of hard liquor per day, two had a flu like syndrome preceding the onset of congestive heart failure, one had drug toxicity preceding the onset of congestive heart failure and no triggering mechanism could be elicited in the remaining two patients.

Discussion

Twenty eight per cent of patients with idiopathic cardiomyopathy, were found to have serum factors reactive to cardiac myofibers at some time during their disease. The appearance or disappearance of these factors did not correlate in this study with exacerbation or remission of disease. If HRF were to be important in the pathogenesis of idiopathic cardiomyopathy we might expect a positive correlation analogous to anti DNA antibody and renal disease in systemic lupus erythematosus but this was not found. Patients with HRF did not have an elevated ASO titer thus making it unlikely that the HRF is a cross reacting antibody to the streptococcus. When present HRF was in low titer in patients with and without exacerbation of disease and did not correlate with severity of disease. The observations are best explained by considering the HRF to be a secondary phenomena produced in response to breakdown of cardiac myofibers after injury to heart muscle.

Twenty eight per cent had a positive latex test but none had a positive SSCAT test. Nineteen per cent had positive thyroid antibody tests. These percentages are significantly higher than controls, none of the patients had evidence of thyroid disease or collagen diseases. None of the patients were over the age of sixty. The results did not correlate with exacerbation and remission of the cardiac disease and are likely a non specific response to the basic disease process.

Twenty five per cent of the patients with idiopathic cardiomyopathy had non specific immunoglobulin elevations. These were generally found in those patients with more

severe disease and increasing congestive heart failure. Several of these patients had congestive hepatomegaly and this might account for some of these elevations. Serum complement as estimated by C3 on immunoelectrophoretic analysis was normal in all sera except one.

There was no correlation between HRF and positive latex tests or positive anti thyroid antibodies. Antinuclear factor was not detected at any time during the course of the disease. This is worth noting because of the high incidence of antinuclear factors in a large number of autoimmune diseases.

Several groups have searched for heart reactive antibody. Robinson and co-workers⁴ noted an 11 per cent incidence of heart reactive antibody and Fletcher and Wenger⁵ found that the frequency was not significantly different from that of a mixed control group. Das and co-workers⁶ using an indirect immunofluorescent method found heart reactive antibody in 6 of 30 patients (17 per cent) with idiopathic cardiomegaly, 7 of 8 patients (88 per cent) with idiopathic hypertrophic subaortic stenosis and in 1 of 43 control subjects.

In a previous study from this center¹ HRF was found in only twelve per cent (12 per cent) of patients studied. Only 12.5 per cent had positive latex tests and one patient had anti thyroid antibody. The higher percentages of positive findings in the present study point out the importance of serial studies as opposed to the single sample study. One of the patients who lacked HRF in the previous study has subsequently developed HRF.

This prospective serial study of the humoral immunologic findings in 32 patients with idiopathic cardiomyopathy like the previous study does not support the concept that immunologic mechanisms are of pathogenetic significance in this disease. Secondary immunologic phenomena are present in IC and are likely to be non specific.

Summary

Thirty two patients with idiopathic cardiomyopathy were followed for 51.6 patient years and evaluated for evidence of immunologic disease. Serologic and immunologic variables were correlated with exacer

bation and remission of the underlying disease. Twenty eight per cent of patients had HRF and 28 per cent had positive latex tests. Nineteen per cent had positive thyroid antibody tests. There was no correlation between positive tests and the clinical course or between positive tests and the severity of the disease. The results suggest that idiopathic cardiomyopathy is a disease in which secondary immunologic mechanisms are present but that they do not play a role in etiology or pathogenesis.

REFERENCES

1. Camp T F, Hess E V, Conway G and Fowler N O. Immunologic findings in idiopathic cardiomyopathy. *Am Heart J* 77:610 1969
2. Fowler N O and Gueron M. Primary myocardial disease. *Circulation* 32:830 1965
3. Fowler N O. Differential diagnosis of cardiomyopathies. *Progr Cardiovasc Dis* 11:113 1971
4. Robinson J, Anderson T and Griebble H. Serologic anomalies in idiopathic myocardial disease (Abstr). *Clin Res* 14:335 1966
5. Fletcher G F and Wenger N K. Auto-immune studies in patients with primary myocardial disease. *Circulation* 37:1032 1968
6. Das S K, Cassidy J T and Petty R E. Antibodies against heart muscle and nuclear constituents in cardiomyopathy. *Am Heart J* 83:159 1972
7. Sanders V and Ritts R E. Ventricular localization of bound gamma globulin in idiopathic disease of the myocardium. *JAMA* 194:171 1965
8. Das S K, Cullen J P, Dodson V N and Cassidy J T. Immunoglobulin binding in cardiomyopathic hearts. *Circulation* 41:612 1971
9. Schur P H and Sandson J. Immunologic factors and clinical activity in systemic lupus erythematosus. *N Engl J Med* 278:533 1968
10. Hess E V, Fink C W, Taranta A and Ziff M. Heart muscle antibodies in rheumatic fever and other diseases. *J Clin Invest* 43:886 1964
11. Kaplan M H, Meyersian M and Kushner I. Immunologic studies of heart tissue IV. Serologic reactions with human heart tissue as revealed by immunofluorescent methods. Iso-immune Wassermann and autoimmune reactions. *J Exp Med* 113:17 1961
12. Kaplan M H. The cross reaction of group A Streptococci with heart tissue and its relation to induced auto-immunity in rheumatic fever. *Bull Rheum Dis* 19:560 1969
13. Kaplan M H and Dallenbach F D. Immunologic studies of heart tissue III. Occurrence of bound gamma globulin in auricular appendages from rheumatic hearts. Relation to certain histopathologic features of rheumatic heart diseases. *J Exp Med* 113:1 1961
14. Ziff M, Brown P, Lospalluto J, Badin J and McEwen C. Agglutination and inhibition by serum globulin in the sensitized sheep cell agglutination reaction in rheumatoid arthritis. *Am J Med* 20:500 1956
15. Balfour B M, Domach D, Rott I M and Couchman K G. Fluorescent antibody studies in human thyroiditis. Autoantibodies to an antigen of the thyroid colloid distinct from thyroglobulin. *Br J Exp Pathol* 42:307 1961

Experimental and laboratory reports

The use of radio-iodinated toluidine blue for myocardial scintigrams*

Edward A Carr Jr MD

Mary Carroll BA

Walter DiGiulio MD

Donald C Blair MD

Ann Arbor and Detroit Mich

Although heart disease is the principal cause of death in many countries and its diagnosis is of major importance methods for direct visualization of the myocardium in intact subjects are few. The roentgenogram of the heart shows the myocardium plus other tissues especially the blood contained in the cardiac cavities as a single shadow. Various techniques show cavities and vessels rather than the myocardium itself. On the basis of the important earlier studies which had demonstrated myocardial uptake of potassium and its analogues it was possible to develop a method for myocardial scintigrams¹ using salts of cesium 131. This method permitted the demonstration of the left ventricular myocardium through the intact chest wall without significant contribution to the image from the cardiac blood pool or adjacent non-cardiac tissues. After original development of the technique in dogs it was extended to humans and the myocardial scintigram has subsequently been shown capable of demonstrating myocar-

dial infarcts^{2,3} aneurysm of the left ventricle^{2,4} cardiomyopathies^{5,6} neoplasm in the ventricular wall¹⁰ and rejection of the transplanted heart.¹¹

However the long biologic half life of cesium creates difficulties not only from the standpoint of radiation dose but also through interference with performance of serial scintigram. As skeletal muscles retain cesium much longer than cardiac muscle^{12,13} the uptake by intercostal muscles interferes with serial myocardial scintigrams during the days that follow injection of a single dose of cesium 131. Moreover its principal emission a 294 KEV x ray poses a considerable problem as a result of its low energy and consequent high absorption in tissues. Therefore although cesium 131 has proved an important radionuclide by permitting the demonstration of the validity of the myocardial scintigram for diagnosis the general clinical usefulness of the myocardial scintigram would be greatly increased by the availability of a more convenient radioactive compound for this

From the Departments of Pharmacology and Internal Medicine, University of Michigan, Ann Arbor, Mich. and St. John Hospital, Detroit, Mich.
*This study was supported partly by U.S. Veterans Public Health Service Grants 5 F11 GM15559 and 5 T01 HE05326.
Received for publication Dec. 27, 1972.
Reprint requests to Edward A. Carr Jr, MD, University of Michigan Medical Center, 4640 Medical Center Ann Arbor, Mich. 48104.
*This work was presented in part at the Fourteenth French Colloquium on Nuclear Medicine, Tours, France, June 6, 1972.

purpose Iodine blue (tolonium chloride) $C_{13}H_{14}N_3SCl$ a phenarazthionium dye was found to concentrate selectively in parathyroid glands of dogs and man after intravenous or intra arterial injection.¹² Toluidine blue has subsequently been found to concentrate heavily in myocardium of dogs after intravenous injection.¹³ Iodotoluidine blue, labeled with iodine 131 or iodine 125 was suggested as a potential radioactive imaging agent for parathyroid scintigrams on the basis of such distribution studies.¹⁴ Moreover a preliminary study¹⁵ showed that unlabeled toluidine blue (hereafter abbreviated TB) was present in decreased concentration in myocardial infarcts as compared to normal myocardium after intravenous injection in dogs. As the possibility of obtaining myocardial scintigrams with a compound of iodine 131 suggested a way of avoiding the problem posed by the weak emission of cesium 131 further investigation of radioactive iodotoluidine blue (hereafter abbreviated as ITB) as a myocardial imaging agent seemed indicated.

But initial studies with ITB were reported as discouraging by other authors as they did not confirm a significant selective uptake of ITB by parathyroid glands¹⁶ or myocardium¹⁷ of rats after intravenous injection of the radioactive compound. It is of interest that the use of a preliminary priming or loading dose of the stable compound TB before injection of the radioactive compound ITB was not reported in any of these negative studies.

We have now demonstrated in both rats and dogs a significant selective uptake of ITB by myocardium when the administration of this radioactive compound is first preceded by a priming dose of the stable compound TB. This uptake by normal myocardium permits demonstration of myocardial infarcts as cold areas of decreased uptake in dogs. We have further demonstrated that the major fraction of administered radioactive iodine given as ITB is excreted with sufficient rapidity to give this compound significant advantage over radioactive cesium as a myocardial imaging agent if serial scintigrams are desired.

Methods

Toluidine blue (TB) USP was dissolved in isotonic saline solution in appropriate concentrations to provide the required dose when given at a rate of 0.02 ml per minute to rats and 1 ml per minute to dogs. Iodinated toluidine blue (ITB) labeled with either iodine 131 or iodine 125 was obtained* in specific activities ranging from 0.3 to 1.2 millicuries per milligram. The dose of ITB given as a bolus intravenously, was 60 microcuries per kilogram of body weight and 20 microcuries per kilogram of body weight for rats and dogs respectively. Rats used in this study weighed from 153 to 410 grams. Dogs weighed from 8 to 17 kilograms.

Studies of the effect of priming doses of stable TB on subsequent distribution of radioactive ITB

RATS Forty nine rats were divided into 7 groups of 7 each and were anesthetized with chloral hydrate. To establish the effect of TB doses ranging from 0 to 10 mg per kilogram of body weight were administered in isotonic saline through a tail vein at a rate of 0.02 ml per minute over various time periods. One group of rats served as controls receiving no TB; this control group received isotonic saline alone prior to the administration of ITB. Immediately following the completion of the infusion of the priming dose all rats were given ITB. The rats were killed 15 minutes after administration of ITB. Tissue samples from all major thoracic and abdominal organs including 4 samples of left ventricular myocardium were taken from each rat. The samples were weighed and counted in a scintillation detector.

DOGS Eight dogs were divided into 4 groups of 2 each. Priming doses of TB ranging from 1 mg per kilogram of body weight to 10 mg per kilogram of body weight were given intravenously at an infusion rate of 1 ml per minute over a period of 45 minutes. Following completion of the infusion of TB, ITB was given intravenously to all dogs. The dogs were killed 15 minutes after administration of ITB. Tissue samples were taken from all the

We are grateful to Gerald Bruno of E. M. Squibb & Sons who generously supplied the radioiodinated toluidine blue.

major thoracic and abdominal organs including six samples of left ventricular myocardium from each dog. The samples were weighed and counted in a scintillation detector.

Studies utilizing a fixed priming dose of stable TB. Following the experiments described above a standard priming dose of 10 mg per kilogram of body weight given intravenously over a period of 45 minutes was selected. Except as otherwise specified all studies described below utilized this priming dose for both rats and dogs. The bolus dose of ITB was given immediately after the conclusion of the infusion of TB.

Retention of radioactivity by myocardium and whole body of dogs after injection of ITB. Eight dogs were divided into four groups of two each. Each dog received a priming dose of TB and a subsequent injection of ITB as described above. These dogs were killed at periods ranging from 15 to 90 minutes after administration of ITB. Tissue samples including six samples from left ventricular myocardium were removed from each dog and were weighed and counted in a scintillation detector.

For an estimation of whole body biologic half life of radioactive iodine when given as ITB three additional dogs were given TB and ITB. The dogs were housed in metabolic cages, excreta were collected and counted daily. The dogs were killed from 7 to 11 days later. A complete set of over 30 tissues was removed from each dog. Appropriate samples were weighed and counted in a scintillation detector. Wherever possible whole organs were weighed; otherwise the weight of whole organs was estimated from a standard reference table.²¹

Two other dogs received TB and ITB. They were studied serially by whole body scintigrams using an Ohio Nuclear Scanner with 3 inch crystal 85 hole collimator and scan minimization recording attachment permitting automatic reduction of the scintigram to one fifth original size. The serial whole body scintigrams were obtained at various times ranging from one half hour to 3 days after injection of ITB.

Myocardial scintigrams. Eleven normal dogs received TB and ITB. Approximately 15 minutes after injection of ITB the dogs were scanned under pentobarbital anes-

thesia in the supine position with the legs extended. In some instances they were subsequently scanned in the left lateral position also. For these scintigrams a Baird Atomic Scanner with 3 inch crystal and 19 hole focusing collimator was used. The scanning speed ranged from 7.5 to 10 inches per minute with the scanner set for sharp contrast (per cent background setting at 80). Further details of the techniques employed have been previously described.²

Experimental myocardial infarcts were induced in 11 dogs by ligation of the anterior descending branch of the left coronary artery under sodium thiopental anesthesia (Except in the two instances noted below none of these dogs had previously been studied among the 11 normals described in the previous paragraph.) After ligation of the coronary artery the chest wall incision was closed and each animal was allowed to recover. Twenty four hours after coronary artery ligation each dog was again anesthetized using pentobarbital. Each dog received 10 mg per kilogram of body weight of TB over a period of 45 minutes followed by 20 microcuries per kilogram of body weight of ITB intravenously as a single bolus dose. Fifteen minutes after the administration of the radioactive compound each dog was killed by an excessive intravenous dose of barbiturate. Myocardial scintigrams were then performed using the same scanning factors employed for the normal dogs that had been scanned alive. After completion of the scintigrams the heart was removed from each killed dog, emptied of all blood and scanned again as an isolated organ. Samples of auricles, right ventricle, lung, liver, rib, muscle, skin and fat were taken for scintillation counting. Three samples of myocardium from areas of anterior left ventricle distal to the site of arterial ligation and three control samples (one from a site proximal to ligation and two from posterior left ventricle areas not dependent upon the ligated artery for their blood supply) were also taken; these were each divided to permit histologic examination as well as scintillation counting. Further details of the techniques employed have been previously described.²² Three of the 11 normal dogs described above were subjected to coronary artery ligation 4 days

purpose Iodine blue (tolonium chloride) $C_{13}H_{16}N_3SCl$ a phenazithonium dye was found to concentrate selectively in parathyroid glands of dogs and man after intravenous or intra arterial injection¹⁴ Toluidine blue has subsequently been found to concentrate heavily in myocardium of dogs after intravenous injection¹⁵ Iodotoluidine blue labeled with iodine 131 or iodine 125 was suggested as a potential radioactive imaging agent for parathyroid scintigrams on the basis of such distribution studies¹⁶ Moreover a preliminary study¹⁷ showed that unlabeled toluidine blue (hereafter abbreviated TB) was present in decreased concentration in myocardial infarcts as compared to normal myocardium after intravenous injection in dogs. As the possibility of obtaining myocardial scintigrams with a compound of iodine 131 suggested a way of avoiding the problem posed by the weak emission of cesium 131 further investigation of radioactive iodotoluidine blue (hereafter abbreviated as ITB) as a myocardial imaging agent seemed indicated.

But initial studies with ITB were reported as discouraging by other authors as they did not confirm a significant selective uptake of ITB by parathyroid glands^{18, 20} or myocardium^{19, 21} of rats after intravenous injection of the radioactive compound. It is of interest that the use of a preliminary priming or loading dose of the stable compound TB before injection of the radioactive compound ITB was not reported in any of these negative studies.

We have now demonstrated in both rats and dogs a significant selective uptake of ITB by myocardium when the administration of this radioactive compound is first preceded by a priming dose of the stable compound TB. This uptake by normal myocardium permits demonstration of myocardial infarcts as cold areas of decreased uptake in dogs. We have further demonstrated that the major fraction of administered radioactive iodine given as ITB is excreted with sufficient rapidity to give this compound significant advantage over radioactive cesium as a myocardial imaging agent if serial scintigrams are desired.

Methods

Toluidine blue (TB) USP was dissolved in isotonic saline solution in appropriate concentrations to provide the required dose when given at a rate of 0.02 ml per minute to rats and 1 ml per minute to dogs. Iodinated toluidine blue (ITB) labeled with either iodine 131 or iodine 125, was obtained* in specific activities ranging from 0.3 to 1.2 millicuries per milligram. The dose of ITB given as a bolus intravenously, was 60 microcuries per kilogram of body weight and 20 microcuries per kilogram of body weight for rats and dogs respectively. Rats used in this study weighed from 133 to 410 grams. Dogs weighed from 8 to 17 kilograms.

Studies of the effect of priming doses of stable TB on subsequent distribution of radioactive ITB

RATS Forty nine rats were divided into 7 groups of 7 each and were anesthetized with chloral hydrate. To establish the effect of TB doses ranging from 0 to 10 mg per kilogram of body weight were administered in isotonic saline through a tail vein at a rate of 0.02 ml per minute over various time periods. One group of rats served as controls receiving no TB; this control group received isotonic saline alone prior to the administration of ITB. Immediately following the completion of the infusion of the priming dose all rats were given ITB. The rats were killed 15 minutes after administration of ITB. Tissue samples from all major thoracic and abdominal organs including 4 samples of left ventricular myocardium were taken from each rat. The samples were weighed and counted in a scintillation detector.

DOGS Eight dogs were divided into 4 groups of 2 each. Priming doses of TB ranging from 1 mg per kilogram of body weight to 10 mg per kilogram of body weight were given intravenously at an infusion rate of 1 ml per minute over a period of 45 minutes. Following completion of the infusion of TB ITB was given intravenously to all dogs. The dogs were killed 15 minutes after administration of ITB. Tissue samples were taken from all the

Table II Radioactivity in tissues of dogs given various priming doses of stable toluidine blue before radio iodinated toluidine blue*

| Priming dose (mg/kg) | Radioactivity $\left(\frac{\% \text{ administered dose}}{\text{Gm tissue}}\right)$ in | | | Ratio of radioactivity | |
|-------------------------|--|--------------|--------------|------------------------|----------------|
| | Left ventricle (LV) | Blood (B) | Liver (L) | $\frac{LV}{B}$ | $\frac{LV}{L}$ |
| 1 | 0.78† | 0.11 | 1.28 | 7.1 | 0.6 |
| 3 | 1.35 | 0.10 | 1.16 | 13.5 | 1.2 |
| 5 | 1.61 | 0.10 | 1.49 | 16.1 | 1.1 |
| 10 | 1.56 | 0.12 | 0.98 | 13.0 | 1.6 |

*See text for details.

†Each value represents the mean of 2 dogs.

cantly ($p < 0.05$) less in rats receiving a priming dose of TB 10 mg per kilogram of body weight over 90 minutes or 10 mg per kilogram of body weight over 180 minutes compared to controls. No other treatment resulted in a significant difference from control values for hepatic uptake. The radioactivity of blood followed the pattern seen in liver with significant ($p < 0.05$) decreases in rats receiving priming doses of 10 mg per kilogram of body weight over 90 minutes or 10 mg per kilogram of body weight over 180 minutes as compared to controls.

Except for liver uptake by non-cardiac tissues adjacent to the heart was not sufficiently high in rats given priming doses of TB to lead to significant interference with the use of ITB as a myocardial imaging agent (see below).

Dogs. As with rats the most important uptake by an organ adjacent to the heart likely to cause interference with the myocardial scintigram was uptake by liver. Table II shows the radioactivity in left ventricle, liver and blood of dogs receiving various priming doses of stable TB. The use of a priming dose increases the ratios radioactivity in left ventricle/radioactivity in blood and radioactivity in left ventricle/radioactivity in liver thus increasing the likelihood of good resolution of the left ventricle by scanning techniques. In dogs

this effect of a priming dose is chiefly by way of increased uptake by left ventricle.

Studies utilizing a fixed priming dose of stable TB

RETENTION OF RADIOACTIVITY BY MYOCARDIUM AND WHOLE BODY OF DOGS AFTER INJECTION OF ITB. The concentration of radioactivity in left ventricle of dogs at various times after injection of ITB is shown in Fig. 1. The high uptake of radioactivity obtained when a priming dose of TB is followed by ITB does not lead to prolonged retention of radioactivity in the myocardium.

From the eight actual measurements obtained during the first 90 minutes after injection the equation shown in Fig. 1 was derived*. The curve of this equation is also shown in the illustration. The uptake at 24 hours after injection calculated from this equation is shown by the dot at the extreme right of Fig. 1. A ninth dog was actually studied at 24 hours and the uptake found in this dog is shown by a bar at the extreme right. The calculated and determined points agree very well. Despite the rapid decay of myocardial radioactivity during the first 90 minutes after injection the ratio of myocardial radioactivity to radioactivity in the blood (not shown in Fig. 1) remained >4 at all times during the

*We acknowledge with thanks the assistance of D. Joh. G. Wagner.

Table I Radioactivity in tissues of rats given various priming doses of stable toluidine blue before radio iodinated toluidine blue*

| Priming dose (mg/kg) and duration of infusion | Radioactivity (% administered dose) in Gm tissue | | |
|---|--|-------|-------|
| | Left ventricle | Liver | Blood |
| Saline control | 0.34% | 0.81 | 0.14 |
| 5 in 15 minutes | 0.49 | 0.43 | 0.10 |
| 5 in 30 minutes | 0.32 | 0.45 | 0.09 |
| 5 in 90 minutes | 0.59 | 0.44 | 0.08 |
| 10 in 45 minutes | 1.20% | 0.67 | 0.10 |
| 10 in 90 minutes | 0.82% | 0.56% | 0.06% |
| 10 in 180 minutes | 0.78 | 0.32% | 0.07% |

*See text for further details

†In this and subsequent tables the administered dose used in the calculation is the number of microcuries given per kilogram of body weight

‡Each value represents the mean of 7 rats

§Differs significantly ($p < 0.05$) from control

after a normal myocardial scintigram had been obtained in the living dogs. Two of these 3 were among the 11 dogs that received a dose of TB and ITB 24 hours after ligation and underwent myocardial scanning after killing. The purpose of sacrificing dogs with myocardial infarcts before scanning was to insure that the subsequent scintillation counting of tissue samples would permit comparable data to be obtained among 11 dogs that had all lived 15 minutes after injection of the radio nuclide. However, one normal dog that subsequently underwent coronary artery ligation and received TB and ITB 24 hours later was not killed. This permitted a scintigram of the infarct to be obtained in a living dog. Although tissue data from this dog could not therefore be compared with data obtained from the 11 dogs killed 15 minutes after ITB injection, this additional dog was killed at the conclusion of the myocardial scanning in order to permit histologic confirmation of infarction.

Results

Studies of the effect of priming doses of stable TB on subsequent distribution of radioactive ITB

RATS: Table I compares the distribution of ITB in rats given various priming doses of TB over various periods of time before injection of ITB. Analysis of variance was

first performed on all uptakes of ITB by left ventricle to determine whether any significant difference existed among the 7 means—to determine whether the use of any priming dose significantly increased uptake over control. After this analysis had demonstrated a significant ($p < 0.05$) difference, further analysis was carried out by Duncan's multiple range test to determine which of the means differed significantly from the control. The latter test demonstrated a significantly higher ($p < 0.05$) uptake of radioactive iodine by left ventricle in rats given 10 mg per kilogram of body weight of TB over a 45 minute period when compared to controls. The respective uptakes were 1.20 ± 0.66 per cent administered dose per gram of tissue and 0.34 ± 0.13 per cent administered dose per gram. Similarly, the uptake by left ventricle in rats receiving 10 mg per kilogram of body weight TB over 90 minutes (0.82 ± 0.35 per cent administered dose per gram) was significantly ($p < 0.05$) higher than uptake by controls. Uptake by left ventricle of rats receiving 10 mg per kilogram of body weight over 180 minutes and uptake by left ventricle of rats receiving priming doses of 5 mg per kilogram of body weight did not differ significantly from controls.

Similar statistical analysis using the multiple range test showed that the uptake of radioactive iodine by liver was significant

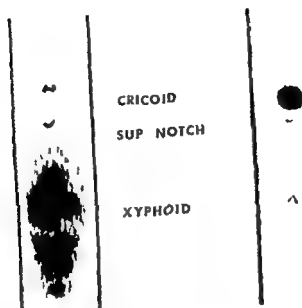


Fig 3 Whole body scintigrams of Dog No. 1 given ITB. Thirty minute after injection of the radioactive compound scanning started at the head and proceeded caudally. The results are shown on the left. The scintigram on the right shows the same dog scanned 3 days later.

calculation of whole body radiation dose using the formulas of Quimby and colleagues² requires an assumption regarding the fate of the 16 per cent which was not found in collected urine, stools and residual carcass. In the best possible case one assumes that this radioactivity was lost very early after administration of the ITB when small errors in collection of the highly radioactive excreta would result in significant losses. In the worst possible case one assumes that the entire 16 per cent was actually in the carcass in addition to the approximately 4 per cent found by analysis. Thus in the best possible case a mean of 71 per cent of total radioactivity was excreted as a short half life component with a biologic half life of 0.63 days. 25 per cent was excreted with a biologic half life of 28 days and 4 per cent remained to be eliminated by physical decay. The respective figures for the worst possible case are 55 per cent, 25 per cent and 20 per cent.

These data in dogs are useful for estimating expected radiation dose in man following the use of ITB. Although species differences obviously may exist and cannot be resolved until studies are conducted in man this preliminary estimation may be of interest. The whole body combined beta

and gamma radiation dose for a 70 kilogram man receiving 1 microcurie per kilogram of body weight of ITB labeled with iodine 131 after an appropriate priming dose of stable TB is estimated on the basis of the above assumptions as 53 millirads in the best possible case and 73 millirads in the worst possible case.

Serial whole body scintigrams of a dog given 10 mg per kilogram of body weight of TB followed by ITB are shown in Fig 3. The scintigrams suggest disappearance of most of the radioactivity from the body within a few days. The results obtained in a second dog subjected to serial scintigrams were very similar.

Myocardial scintigrams. Figs 4 and 5 show anterior views of normal and infarcted myocardium respectively. Although our tissue studies have repeatedly shown that uptake by normal right ventricle and atria is comparable to uptake by normal left ventricle the myocardial scintigram results almost exclusively from radiation emanating from the left ventricle as a consequence of the preponderant mass of the latter. Figs 6 and 7 show lateral views of normal and infarcted left ventricle respectively. These four scintigrams were obtained in 4 different dogs. Fig 8 however

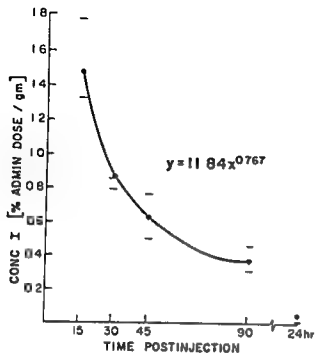


Fig 1 Concentration of radioactive iodine in the left ventricle of dogs at various times after intravenous injection of ITB. Horizontal line (—) indicates measured value. The dot at 24 hours represents a calculated value. See text for details.

first 90 minutes. The average of this ratio for the eight points during that time was 10.9.

The retention of radioactivity in dogs as calculated from daily urine and stool collections is shown in Fig 2. At the time of death, the total radioactivity remaining in the carcass of dog A (killed at 7 days), dog B (killed at 10 days), and dog C (killed at 11 days) was 3.28, 6.1, and 3.84 per cent of administered dose, respectively. By the usual curve stripping method, the retention curve of radioactivity was separable into two components as shown in Fig 2. The first component had a half life of 0.6, 0.6, and 0.7 days, respectively, in the 3 dogs (mean, 0.63 days). The second component had a half life of 24, 32, and 28 days, respectively, in the 3 dogs (mean 28 days). Extrapolation of the second component back to the origin indicated that 24 per cent, 31 per cent, and 20 per cent of the total dose was retained in the 3 dogs, respectively, with a mean biologic half life of 28 days. In estimating whole body radiation dose, we have made the conservative assumption that the residual in the carcass would not have been excreted and would have been lost only through physical decay.

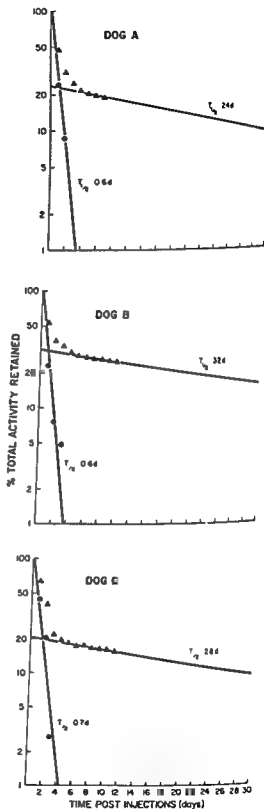


Fig 2 Whole body retention of radioactivity by dogs given ITB.

As it was possible to account for a mean of 84 per cent of the administered dose of radioactive iodine through the total collected in urine and stools and the analysis of residual radioactivity in the carcass



Fig. 6 Scintigram of a normal dog heart (Dog No. 4) Lateral view. The priming dose of TB was 5 mg per kilogram of body weight in this dog.



Fig. 7 Scintigram of Dog No. 5 showing a myocardial infarct. Lateral view.

injection of ITB the mean ratio at time of death in these studies was approximately 12 to 1. The scintigram after administration of ITB in these experiments represents primarily radioactivity from the myocardium with little contribution from the low level of activity contained in the blood

in the heart cavities. This was repeatedly confirmed by scintigrams of the isolated hearts removed from the dog and emptied of all blood. The emptied hearts gave scintigrams essentially the same as hearts which contained blood in their cavities showing normal left ventricle in the normal



Fig 4 Myocardial scintigram of a normal dog heart (Dog No 2) Anterior view For this and all subsequent figures a priming dose of stable TB was given intravenously over a 45 minute period followed by a bolus injection of radioactive ITB Scintigram began 15 minutes after the latter injection Except where otherwise specified the dose of TB was always 10 mg per kilogram of body weight and each dog was killed at the beginning of the scintigram



Fig 5 Scintigram of Dog No 3 showing a myocardial infarct Anterior view Infarcts shown in this and all subsequent figure were confirmed by histologic examination

shows anterior scintigrams obtained in the same dog before and one day after experimental myocardial infarction Fig 9 shows lateral scintigrams obtained in still another dog before and one day after experimental

infarction Although the ratio concentration of radioactivity in normal left ventricle/concentration of radioactivity in whole blood varies among individual dogs and varies to some degree with time after



Fig 10 Myocardial scintigrams of isolated hearts. Left normal heart. Right heart showing myocardial infarct at the apex. A normal dog and a dog with a myocardial infarct respectively each received TB and ITB. The respective TB doses were 1 mg per kilogram of body weight and 10 mg per kilogram of body weight. Fifteen minutes after injection of ITB the dogs were killed and their hearts were removed. After all blood had been emptied from the heart cavities as previously described the hearts were scanned. The actual outlines of the heart are projected on the scan. The density represents left ventricular muscle.

scintigrams may be obtained by repeated doses of ITB even if labeled with iodine 131.

Although the rapid disappearance of radioactivity from the myocardium after administration of ITB requires that scintigrams be performed promptly after administration of ITB, this has not caused significant problems. The radioactivity of the normal left ventricle remains much higher than the radioactivity of infarcted areas or of blood even several hours after administration of ITB—i.e. there is a comparable decline of radioactivity in all three types of tissue. Therefore the ratio of radioactivity in target tissue to that in non target tissue, the most important factor in permitting a good scintigram, remains very favorable for a considerable period of time.

Thus the use of ITB offers the possibility of myocardial scintigrams that do not suffer from either of the two disadvantages of cesium 131: weak emission and long biologic half life. Cesium 131 has already been shown to be effective in man in demonstrating myocardial infarcts and other myocardial lesions. The possibility of obtaining myocardial scintigrams with the more advantageous radioactive compound may significantly improve diagnosis of

myocardial infarction in the future and may make it possible to evaluate more precisely therapeutic regimens in myocardial infarction. It is of interest that the mean uptake of infarcted myocardium is about 15 per cent the uptake by normal myocardium, whether one uses labeled toluidine blue or radioactive cesium.

To our knowledge the closest previous approach to demonstration of myocardial infarcts using a compound labeled with radioactive iodine was that reported by Gunton and associates²⁴ who reported that oleic acid labeled with iodine 131 could be complexed to human serum albumin and utilized in a technique which, according to the authors' report, just approached the limits of definition of infarction when used for scintigrams in acute cases. As noted above, iodine 131 is likely to be a very satisfactory label for ITB. We have not thus far attempted to use iodine 123. However, despite the short physical half life of iodine 123, its emission has a particularly favorable energy (159 keV) for use with a radiation camera. Therefore the eventual possibility of using iodine 123 should not be entirely discounted, especially if one wishes to use a radiation camera rather than a scanner.



Fig 8 Myocardial scintigrams of Dog No 6 before and after myocardial infarction. Anterior view. The scan on the left was obtained one day before infarction. The scan on the right was obtained 4 days after infarction. Both scintigrams were performed in the living dog.



Fig 9 Myocardial scintigrams of Dog No 7 before and after myocardial infarction. Lateral view. The dog was studied in the same manner as the dog shown in Fig 8. This dog was alive during the performance of the scan on the left and was killed just before performance of the scan on the right.

dogs and cold areas of decreased uptake in dogs with myocardial infarcts (Fig 10).

Table III summarizes the data obtained in 11 dogs with myocardial infarctions. Actual necrosis of tissue was histologically confirmed in every instance. The mean ratio concentration of ITB in infarcts/concentration in normal left ventricle was 0.18 ± 0.31 . The difference in uptake by normal and infarcted tissue was significant ($p < 0.005$) by the t test.

Discussion

Both iodine 131 and iodine 125 were used as labels in these experiments in animals. The availability of both labels for ITB offers in itself the possibility of serial

scintigrams within a short interval of time as one may easily detect iodine 131 while discriminating against iodine 125. The usefulness of both of these isotopes in laboratory animals should not obscure the fact that iodine 131 with its more penetrating emission is likely to be the better isotope in man. The emission of iodine 125 would not represent a significant advantage over that of cesium 131, while iodine 131 has a significant advantage in this respect. The rapid disappearance of radioactivity through excretion is an additional advantage permitting serial scintigrams without excessive radiation dose. The rapid disappearance of radioactivity from the myocardium strongly suggests that serial

Therefore these data are recorded only to suggest that the requirement of a priming dose of stable toluidine blue need not pose an insuperable barrier to the use of ITB as a myocardial scanning agent. Although the animal studies reported here are promising the authors do not of course advocate that any initial trial in humans will require 10 mg per kilogram of body weight as a starting dose regardless of the rate of infusion. The same general rules that govern early human trials of therapeutic compounds must govern trials of diagnostic agents—the initial approach must be cautious utilizing low doses. Moreover studies in normal subjects should precede studies in patients.

REFERENCES

- 1 Love W D, Romney R B and Burch G E. A comparison of the distribution of potassium and exchangeable rubidium in the organs of the dog using rubidium 86. *Can Res* 2:112 1954
- 2 Carr E A Jr, Goss G, Shaw J and Krontz B. The direct diagnosis of myocardial infarction by photoscanning after administration of cesium 131. *Am Heart J* 68:627 1964
- 3 Carr E A Jr. The development of myocardial scanning with special reference to the use of cesium 131 in man. In Quinn J L. (ed) *Proceedings of the Winston Salem Scanning Symposium Scintillation Scanning in Clinical Medicine*. Philadelphia 1964 W B Saunders Company p 93
- 4 Goodrich J K, Store H L, Harris C C and Hill R. Clinical applications of low energy high transmission collimator. *J Nucl Med* 6:109 1965
- 5 McGeehan J T, Rodriguez Antune A and Lewis R C. Cesium 131 photoscan Aid in the diagnosis of myocardial infarction. *JAMA* 204:185 1968
- 6 Brochier M, Haniel T, Garner G, Archambault D and Raymond R. La scintigraphie du myocarde et de ses lésions. *Arch Mal Coeur* 64:921 1971
- 7 Haniel T, Garner C, Itu R and Brochier M. La scintigraphie cardiaque. *J Biol Med Nucl* 24:17 1973
- 8 Inoh T. Study on the idiopathic cardiomyopathy. Diagnostic value of the apexcardiogram with scintiscanning of the heart and plasma hydroxyproline. *Jap Circ J* 35:731 1971
- 9 Riquet W and Merchie G. La scintigraphie myocardique au césium 131. *Rev Med Liege* 27:333 1972
- 10 Quasle M A and Wilson W J. Detection of cardiac tumor by rectilinear imaging with cesium. *J Nucl Med* 14:605 1970
- 11 Carr E A Jr, Kahn D R, Carro M, Oberman H A and Dufek J H. The uptake

- of radioactive cesium by transplanted human and animal hearts: theoretical and practical significance. *Int J Clin Pharmacol Ther Toxicol* 4:72 1970
- 12 Ballou J E and Thompson R C. Metabolism of Cesium 137 in the rat. *Health Phys* 1:85 1958
- 13 Carr E A Jr. Pharmacology of radioactive cesium salts. In *Proceedings of the Oak Ridge Inst. of Nuclear Studies Symposium on Radioactive Pharmaceuticals*. U.S. Atomic Energy Commission Division of Technical Information 1966 p 619
- 14 Kopper I J and Moe R E. Demonstration of parathyroid during surgery in dogs with preliminary reports of results of some clinical cases. *Surgery* 59:1101 1966
- 15 DiGiulio W and Lindenauer S M. Use of technetium chloride in localization of parathyroid tissue. *JAMA* 214:2302 1970
- 16 Kagan G S and DiGiulio W. Potential value of toluidine blue analogs as parathyroid scanning agents. *J Nucl Med* 9:643 1968
- 17 Kagan G S. Potential value of toluidine blue analogs as scintiscanning agent for myocardial infarcts. *J Nucl Med* 10:413 1969
- 18 Archer E G, Potchen E J, Studer R and Siegel B. Distribution of ¹²⁵I toluidine blue in the rat. *J Nucl Med* 10:386 1969
- 19 Larose J H, Whitaker R H and Reba R C. Radioiodinated toluidine blue: an unsatisfactory scanning agent. *J Nucl Med* 11:731 1970
- 20 Archer E G, Potchen E J, Studer R and Siegel B. Tissue distribution of ¹²⁵I toluidine blue in the rat. *J Nucl Med* 13:85 1972
- 21 Spector W S. (ed) *Handbook of biological data*. Philadelphia 1966 W B Saunders Company p 163
- 22 Carr E A Jr, Bernales W R, Patno M E, Bartlett J D Jr and West A V. The detection of experimental myocardial infarcts by photoscanning. *Am Heart J* 64:650 1962
- 23 Quimby E A and Feitelberg S. Radioactive isotopes in medicine and biology. Philadelphia 1963 Lea & Febiger Publishers chapter 11
- 24 Gaulton R W, Evans J R, Baker R C, Spears J C and Beaman D S. Demonstration of myocardial infarction by photoscans of the heart in man. *Am J Cardiol* 16:382 1966
- 25 Kiese M, Lorcher W, Weiser J and Zuber A. Comparative studies on the effects of toluidine blue and methylene blue on the reduction of ferric haemoglobin in man and dog. *Fur J Clin Pharmacol* 4:115 1972
- 26 Allen J G, Grossman E, Elghamry R M, Moulder P V, McKeen C L, Jacobson L O, Pierce M, Smith F R and Crosbie J. Abnormal bleeding. Response to treatment with toluidine blue and, sodium sulfate. *JAMA* 139:1731 1949
- 27 Eastwood G L. ECG abnormalities associated with the barium enema. *JAMA* 219:19 1972

Table III Radioactivity in hearts of dogs with myocardial infarcts*

| Dog | Uptake† by | | Uptake by normal LV Uptake by infarcted LV | Radioactivity in normal LV radioactivity in blood |
|-------------|------------------|---------------------|---|--|
| | Normal LV tissue | Infarcted LV tissue | | |
| A‡ | 0.88 | 0.11 | 0.13 | 13.62 |
| B‡ | 0.98 | 0.10 | 0.10 | 13.07 |
| C‡ | 0.30 | 0.015 | 0.05 | 10.24 |
| D | 0.28 | 0.31 | 1.11 | 9.83 |
| E | 1.48 | 0.07 | 0.05 | 9.16 |
| F | 1.22 | 0.22 | 0.18 | 19.6† |
| G | 0.87 | 0.14 | 0.16 | 9.34 |
| H | 0.93 | 0.075 | 0.08 | 7.15 |
| I | 1.36 | 0.06 | 0.04 | 8.80 |
| J | 0.51 | 0.006 | 0.01 | 18.92 |
| K | 0.29 | 0.02 | 0.07 | 8.29 |
| Mean ± S.D. | 0.83 ± 0.43 | 0.10 ± 0.09‡ | 0.18 ± 0.31 | 11.64 ± 4.23 |

*Each dog received 100 mg per kilogram of body weight stable toluidine blue over 45 minutes preceding the bolus dose of radio-iodinated toluidine blue. Each dog was killed 15 minutes after the latter dose. See text for further details.

†Per cent administered dose/Gm

‡Differs significantly ($p < 0.005$) from normal left ventricular tissue

§Not the same dogs as dogs A, B, and C of Fig. 2

The mechanism whereby a priming dose of TB first suggested by one of the authors (W. DiG.) influences the distribution of ITB is at present unknown. Both the rate of administration and the total dose of stable TB are significant as suggested by the studies in rats. The use of a priming dose of TB does not increase the concentration of radionuclide in important non-target tissues (liver and blood). Indeed, the data in rats suggest lower radioactivity in these non-target tissues as a result of a priming dose of TB. It should be noted that the use of a priming dose does not increase the uptake of radionuclide by various other non-cardiac tissues after administration of ITB (e.g., muscle, lung, etc.) but data on liver and blood have been presented here as they represent particularly important non-target tissues in the myocardial scintigram.

One may postulate two types of binding site for the phenazathionium nucleus such that one of the sites has a lower capacity but higher affinity than the other. Saturation of the first site—e.g., in liver and blood—by a priming dose of TB might then make more ITB available for the second site in the myocardium. Such a mechanism is at present entirely speculative although the

reported electrocardiographic changes after an intravenous injection of TB (see below) may be pertinent here.

Toluidine blue has been used intravenously in man^{14, 15, 16} in the past and has not proved to be a highly toxic compound. Kiese and associates¹⁴ gave 4 mg per kilogram of body weight of TB intravenously to humans without ill effect. Doses of 6 mg per kilogram of body weight have been given safely to humans intravenously¹⁵; these doses were given over a period of 2 hours. Nevertheless, rapid infusion of a sufficiently high dose may result in electrocardiographic changes. Whereas eight patients studied by DiGiulio and Lindenauer¹⁶ received 5 mg per kilogram of body weight of TB intravenously over periods ranging from 45 to 135 minutes without reported electrocardiographic abnormalities, one patient who received 3.5 mg per kilogram of body weight in 30 minutes did show transient depression of the ST segment.

The development of cardiac arrhythmias after diagnostic procedures is by no means unknown and even the barium enema is not innocuous in this regard¹⁷ but particular caution is clearly needed to avoid precipitating cardiac arrhythmias in patients already suffering from myocardial infarcts.

striking that our curiosity was aroused. This study has enabled us to define its physiological and clinical significance.

Methods

We used a capacitance Brecht and Bouche transducer having a frequency response flat to 200 Hz.² The transducer was attached to a short rod which projected 1.5 cm beyond a metal circle which surrounded it. The tip of this rod placed over a point where maximal pulsation in the carotid had been palpated was pushed down until the circle was in contact with the skin and then it was held firmly in place with a strap around the neck. A standard differentiating circuit and two channels of a Sanborn Twin beam recorder were used to record the pulse and its derivative simultaneously. The methods of calibrating the pulse derivative (PD) and of measuring the maximum rate of pressure rise (ID max) have been described.⁴

Such records were always taken after the supine subjects had rested comfortably for at least 15 minutes. Several positions of the transducer were tried in each subject that producing the largest record—i.e. the record requiring the least amplification being preferred. ULF force ballistocardiograms taken simultaneously with the pulse records play a very minor part in this study.

When right and left heart catheterizations were performed they were done a day or two before or after the pulse records were taken. Pressures obtained at catheterization were inscribed by an Electronics for Medicine DR12 polybeam photographic recorder using a Stratham P23 DB transducer. The transaortic aortic gradient was measured during pullback from the left ventricle the systolic mean gradient being recorded. We used the conventional 20 mm Hg as the upper limit of normal for the aortic valve gradient.

Subjects

The patients studied were adults who had or were suspected of having serious cardiac disease. All were inpatients in the University Hospital and all were ambulatory.

One hundred-eighty patients were subjected to cardiac catheterization. Since the

composition of this group was biased by the ordinary criteria for selecting patients for this test a larger and a more representative group of cardiac patients was secured by adding the last 169 cardiac patients studied who had not been catheterized records of their pulse derivatives having been taken in conjunction with ballistocardiographic studies. The final hospital diagnosis used to classify each case was made by a doctor without knowledge of our findings in the pulse.

Results

What we have called the carotid high frequency abnormality illustrated in Figs 1 and 2 consists of a brief burst of high frequency distortion superimposed on the basic pattern of the pulse derivative for part of the cardiac cycle. Usually of 15 Hz or faster the burst starts at or shortly after the main wave's peak and usually lasts from 0.1 to 0.4 second. By its regular position at this part of the cardiac cycle the abnormality can be easily distinguished from electrical noise.

As shown in Figs 1 and 2 when the abnormality was present and the rhythm regular the general pattern was repeated with every heart beat but there was often some variation in the HF details of the burst. Such small differences are probably due to artefacts for slight changes in the exact positioning of the sensor on the artery and in the pressure applied often caused similar changes in the record probably by altering the transmission of the HF vibrations from arterial lumen to sensor. No similar bursts of HF distortion have ever been encountered in pulse derivatives secured on healthy persons so the abnormality can be recognized at a glance (Fig 1). Duplicate runs made on the same day or on different days give excellent agreement of the general pattern of the abnormality despite some differences in the HF details.

Very conspicuous in the pulse derivative the HF abnormality can sometimes be recognized in the conventional pulse but only if one examines the record very carefully (Fig 1) in the majority of cases it would certainly be overlooked.

Table I was compiled from data secured from the 180 patients catheterized. There

Disturbed blood flow in the carotid artery Its physiological and clinical significance

Isaac Starr MD*

Christophe Ambrosi MD*

Joel H Manchester MD**

James C Shelburne MD**

Philadelphia Pa

Elasticity in the rubber membranes rubber tubing and air transmission systems used in the early clinical studies of the pulse prevented the proper recording of its high frequency (HF) components so one cannot expect to find evidence of HF abnormalities in this literature. We know of no mention of such abnormalities before 1955 when Smith¹ after improving his apparatus by greatly diminishing the size of his air transmission system described an HF abnormality in the carotid pulse of a case of aortic stenosis. Smith also performed a very illuminating experiment on this patient after he increased the size of his air transmission system until it resembled the apparatus used in early work on the pulse, the HF abnormality disappeared and was replaced by the 'bisferiens' type of pulse so often described as characteristic of aortic stenosis in the older literature.²

Soon after Smith's study the use of rigid apparatus in conjunction with electrical

methods of transmission and recording provided records in which the HF components of the pulse were much better recorded and Smith's finding was soon confirmed in several laboratories.³⁻⁷

In this laboratory increasing interest in the cardiac forces and in incoordination of the cardiac contraction led to continued efforts to improve our knowledge of the HF components of the pulse. Thus since 1959 the pulse derivative has been recorded routinely to bring out HF information and, in 1964 the sensor was shifted from the brachial to the carotid artery to minimize the loss of HF information which occurs as the wave travels down the artery. Since that time pulse records containing far more HF information than has been recorded in most other clinical studies of the pulse have been secured in over 1 000 subjects both patients and healthy persons.

Certain of these patients exhibited an HF abnormality of the carotid pulse so

From the Department of Therapeutic Research and the Department of Medicine, Cardiac and Pulmonary Division, School of Medicine, University of Pennsylvania, Philadelphia, Pa.

Received for publication Dec 27 1972

Reprint requests to Dr Isaac Starr, University of Pennsylvania, 851 Gates Memorial Pavilion, Philadelphia, Pa 19104.

*Drs Starr and Ambrosi were supported by grant No. HL 625 CVB from the United States Public Health Service, National Institutes of Health.

**Drs Manchester and Shelburne were supported by grant No. HL 8805 from the United States Public Health Service, National Institutes of Health.

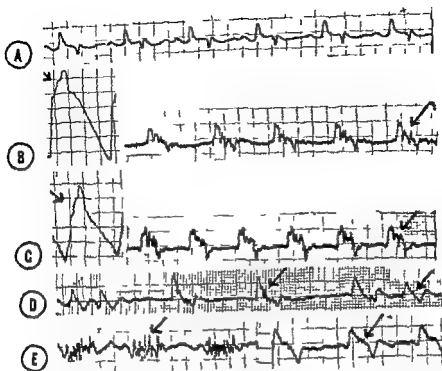


Fig 1 A through E High frequency abnormalities in the carotid pulse. Arrows point to features of interest. A Normal record for comparison on PD max = 680 mm Hg per second. B Enlarged carotid pulse (left) and pulse derivative (right). Patient K, V, age 18, gradient 37 mm Hg. Congenital aortic stenosis, aortic insufficiency Grade 2. Note conspicuous HF abnormality in PD and that they can also be seen in the enlarged conventional pulse. PD max = 477 mm Hg per second. C Same records of patient F, U, age 54, gradient 57 mm Hg. Aortic stenosis, aortic insufficiency Grade 3. Note conspicuous abnormality in PD; it is present but almost unrecognizable in the enlarged conventional carotid pulse. PD max = 324 mm Hg per second. D Carotid PD of patient T, R, age 73, RHD, severe aortic and mitral regurgitation, atrial fibrillation, gradient variable, minimum 10 mm Hg. Note HF abnormality in large beat, its absence from smaller B. Carotid PD (left) and brachial PD (right) of D, C, age 33. Aortic stenosis and aortic insufficiency Grade 4. Gradient 80 mm Hg. PD max = 248 mm Hg per second. Note maximum HF abnormality in carotid PD, its almost complete disappearance by the time the brachial was reached.

physical facts both a distorted aortic lumen and a flow velocity which exceeds a critical level are needed to produce the bursts of disturbed flow we detect in our patients.

Classic physical experiments indicate that similar factors are concerned in the genesis of disturbed flows in tubes.⁹ Roughness of the wall and irregularities in the shape of the tubes such as narrowing and widening will promote the appearance of turbulence whenever flow velocity exceeds a critical level. Nevertheless turbulence as usually defined today is a phenomenon of much higher frequency than our methods of recording the pulse would detect.¹⁰ Thus while evidence secured by Doppler techniques¹⁰ indicates that blood turbulence does indeed accompany some cases of

aortic stenosis our studies provide no evidence for or against its presence in our cases.

So our data suggest that the carotid high frequency abnormality found in our patients is due to a disturbance of flow profile in which normal streamlined flow is interrupted by eddies, vortices and jets. This abnormality should not be called turbulence but if sufficiently intense it would probably be accompanied by turbulence. Such a physiologic abnormality would like turbulence handicap the heart by increasing the impedance to ejection. So we have demonstrated that a physiological abnormality of blood flow which handicaps cardiac performance is present in most cases of aortic stenosis.

Relation of disturbed aortic flow to ab

Table I Primary cardiac diagnoses made on 180 patients catheterized and their relation to the carotid high frequency abnormality

| Diagnosis | Patients with the HF abnormality (No) | Patients without the HF abnormality (No) |
|---|---------------------------------------|--|
| Calcific aortic stenosis | 5 | 11 |
| Aortic stenosis (non calcific) | 2 | 2 |
| Subaortic stenosis | 1 | 0 |
| Aortic stenosis and insufficiency | 15 | 1 |
| Aortic insufficiency without stenosis | 11 | 13 |
| Mitral valve disease | 7 | 27 |
| Coronary artery disease | 1 | 57 |
| Congenital septal defect or patent ductus | 0 | 12 |
| Cardiomyopathy | 0 | 6 |
| Hypertensive heart disease | 0 | 5 |
| Congenital pulmonary stenosis | 0 | 3 |
| Other cardiac lesions | 0 | 3 |
| No cardiac diagnosis | 0 | 9 |

is a highly significant relationship between the presence and absence of the carotid HF abnormality and the normality and abnormality of the gradient χ^2 square = 63

The relations between the presence and absence of the HF abnormalities and the final clinical diagnoses in the group catheterized are shown in Table I. In the enlarged group of 349 patients the HF abnormality was present in 89 per cent of 35 cases diagnosed aortic stenosis in 48 per cent of 29 cases diagnosed aortic regurgitation with out stenosis and in less than 1 per cent of 285 cases diagnosed cardiac disease without aortic valve involvement. It has not been seen in any healthy person.

Obviously there is a highly significant relation between the carotid HF abnormality and aortic valve disease.

Discussion

Relation to the thrill The genesis of the HF abnormality demonstrated in our patients is surely related to that of the thrill palpable in so many cases of aortic stenosis but our modern methods detect the abnormality in many cases in which no thrill was discovered at routine examination. Evi-

dently we have a more delicate test than palpation provides.

Genesis of the carotid HF abnormality An array of clinical evidence indicates that an anatomical factor associated with the aortic valve is of prime importance in the genesis of this abnormality. It was very seldom encountered in patients without manifest aortic valve disease. In all four cases in which an aortic valve prosthesis was inserted into a previously stenotic orifice, the HF abnormality disappeared (Fig 2) or was greatly ameliorated although lower frequency abnormalities often remained in the pulse record.

But our evidence also points to the importance of a physiological factor in its causation. In all cases the HF distortion occurred in only one part of the cardiac cycle and after the time of the normal peak of the pulse derivative when ejection velocity would be at its highest (Figs 1 and 2). In some cases in which the heart beats differed in strength (Fig 1), the abnormality was present whenever the pulse was large and absent when it was small. In one case (Fig 3) the HF abnormality disappeared after coronary bypass surgery, to reappear after the heart had recovered from the immediate effects of the operation. Such differences are certainly not due to structural changes in an aortic lesion; a physiological factor such as differences in flow velocity must be invoked to account for the findings.

One patient later demonstrated an operation to have a tight aortic stenosis never exhibited the carotid HF abnormality. Before operation his PD max was one of the smallest encountered in this study and his cardiac output (dye method) was calculated to be only 1.4 liters/min/m². Apparently his blood velocity was too small to generate the HF abnormality despite the presence of the aortic lesion.

Several cases of wide open aortic regurgitation whose ejection acceleration must have been very great to account for their large force ballistocardiograms and huge PD max's (Fig 3) showed no trace of the carotid HF abnormality. Evidently neither a tight stenosis of itself nor a high ejection velocity of itself is sufficient to produce the abnormality. So our evidence leads to a theory which is consistent with well known

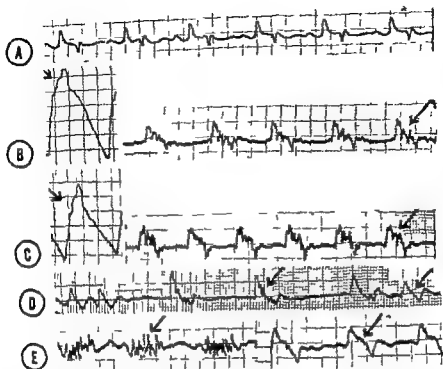


Fig 1 A through E High frequency abnormalities in the carotid pulse. Arrows point to features of interest. A A normal record for comparison. PD max = 680 mm Hg per second. B Enlarged carotid pulse (left) and pulse derivative (right). Patient K. N. age 18. Gradient 37 mm Hg. Congenital aortic stenosis, aortic insufficiency, Grade 2. Note conspicuous HF abnormality in B and that they can also be seen in the enlarged conventional pulse. PD max = 477 mm Hg per second. C Same records of patient F. U. age 54. Gradient 37 mm Hg. Aortic stenosis is aortic in efficiency, Grade 3. Note conspicuous abnormality in PD. It is present but almost unrecognizable in the enlarged conventional carotid pulse. PD max = 324 mm Hg per second. D Carotid PD of patient T. R. age 23. RHD, severe aortic and mitral regurgitation, atrial fibrillation, gradient variable, minimum 10 mm Hg. Note HF abnormality in large beats, its absence from smaller. E Carotid PD (left) and brachial PD (right) of D. C. age 33. Aortic stenosis and aortic insufficiency, Grade 4. Gradient 60 mm Hg. PD max = 248 mm Hg per second. Note maximum HF abnormality in carotid PD, its almost complete disappearance by the time the brachial was reached.

physical facts both a distorted aortic lumen and a flow velocity which exceeds a critical level are needed to produce the bursts of disturbed flow we detect in our patients.

Classic physical experiments indicate that similar factors are concerned in the genesis of disturbed flows in tubes.⁸ Roughness of the wall and irregularities in the shape of the tubes, such as narrowing and widening, will promote the appearance of turbulence whenever flow velocity exceeds a critical level. Nevertheless, turbulence as usually defined today is a phenomenon of much higher frequency than our methods of recording the pulse would detect.⁹ Thus, while evidence secured by Doppler techniques¹⁰ indicates that blood turbulence does indeed accompany some cases of

aortic stenosis, our studies provide no evidence for or against its presence in our cases.

So our data suggest that the carotid high frequency abnormality found in our patients is due to a disturbance of flow profile in which normal streamlined flow is interrupted by eddies, vortices, and jets. This abnormality should not be called turbulence, but if sufficiently intense it would probably be accompanied by turbulence. Such a physiologic abnormality would like turbulence handicap the heart by increasing the impedance to ejection. So we have demonstrated that a physiological abnormality of blood flow, which handicaps cardiac performance, is present in most cases of aortic stenosis.

Relation of disturbed aortic flow to ab



Fig. 2. A through D. Effect of the insertion of a prosthetic caged ball valve into a stenotic aortic orifice. A and B. Cirod PD of Δ U. The 68. Luteic aortic teno: and in efficiency. A Before operation gradient 47 mm Hg. PD max = 421 mm Hg per second. B After operation PD max = 817. Note the appearance of motion but not all of the cirod HI abnormality. C and D. Same records on Δ U. The 58. Aortic teno: aortic in efficiency. Hg/cc. C Record taken before operation when gradient = 50 mm Hg and PD max = 715 mm Hg/cc. D Record taken after operation when PD max was 777 mm Hg/cc. Note that the HI di tortion di appear after operation but some low frequency di tortion remain.

normal aortic valve gradients. The disturbed flow indicated by the HI abnormality in the pulse is accompanied by an abnormal gradient in almost all our cases. Theoretically the two are related in several ways.

Velocity being constant, abnormal pressure drops across the aortic orifice might conceivably be due to impedance caused by narrowing alone or by disturbed flow alone. In the great majority of our cases the increased gradient was probably due to both factors, how much each contributed being unknown. Impedance due to disturbed flow alone may be larger than most doctors realize. Schultz and colleagues¹⁴ perfused normal and arteriosclerotic abdominal aortas secured at necropsy with a fluid of the same viscosity as blood in normal vessels. The gradient at a velocity of 40 cc per second was 2 to 3 mm Hg. But in one sclerotic aorta with a maximum narrowing of only 6 per cent, a gradient of 12 mm Hg was found. Since this amount of narrowing would not cause a detectable gradient in a normal vessel, this fourfold increase of gradient must be attributed to the impedance of a

flow disturbed by roughening and distortion of the vessel walls. Obviously the disturbed flow pattern demonstrated in our patients may be an important factor in their disability.

From the physiological viewpoint both gradient and flow disturbance are influenced by blood velocity, but in a different way. Venturi's principle describes the relation of narrowing to the gradient; in this formula velocity appears as a continuous function. In equations describing the relations of turbulence and disturbed flows, velocity is a step function—i.e., the phenomenon appears when velocity exceeds a critical level. So it is easy to see why an abnormal gradient and a disturbed flow are usually associated in our patients and also that they need not always be associated. Our clinical data on the latter situation are of interest.

Discussion of unusual findings. In 17 cases the HI abnormality was present but the pressure gradient across the aortic valve was not abnormally increased. Most of these were cases of aortic valve disease in which regurgitation predominated. Ev-

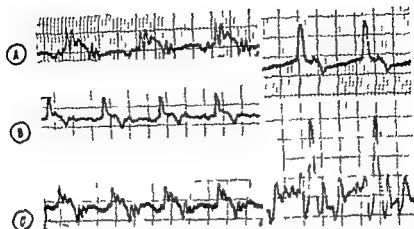


Fig 3 Left panel A B and C Factor in the presence or absence of the carotid HF abnormality Carotid pulse derivative of B T a e 44 diagnosed coronary heart disease A Before operation note HF abnormalities B Twelve days after coronary bypass operation note disappearance of HF abnormalities C Two month post-operative note return of HF abnormality PD max was 300 276 and 300 mm Hg per second in the three records Right panel Carotid pulse derivative over force BCG of V R age 23 wide open aortic regurgitation PD max = 4,500 mm Hg per second which is record breaking Calibration of force BCG 13 small div is on = 275 (10%) dynes So this record is also huge The aortic valve gradient was normal Note that in the presence of what must have been a very great blood velocity HF abnormalities in the carotid PD are absent

dently a roughening without narrowing of the orifice may produce a disturbed flow causing too little impedance to produce gradient abnormalities.

In five cases the gradient was found to be abnormally increased but no disturbed flow was detected. Such findings might have been due to an artefact for which we were on the lookout: should any patient be more excited at catheterization than when the pulse derivative was taken such findings might well result. However the pulse rates recorded at the two tests were so similar in each of these five patients that it is hard to believe that differences in excitement were the cause of our findings. It is easier to believe that the increased impedance in these cases was due to aortic narrowing alone, the lesion having so little roughness that a disturbed flow did not accompany it.

The possibility that disturbed flow in the carotid might be created without an aortic lesion should also be discussed for we have five cases in which the HF abnormality was present without any clinical evidence of aortic valve disease. All these were cases of advanced rheumatic heart disease with several mitral valvulitis. To account for murmurs heard in the aortic area when that valve is normal it has been suggested

that when blood regurgitates through a deficient mitral valve in sufficient amount the jet strikes the atrial wall adjacent to the aortic arch such a jet might not only produce an audible murmur but might also set up vibrations in the aortic blood which our sensor would detect. While one should not forget that an aortic lesion so common in advanced cases of RHD might have been present in these five cases though undiagnosed those doing the catheterization were convinced that this valve was normal.

It is also conceivable that small elements of a myocardium which had lost the coordination of its contraction might beat such a rapid tattoo on the ventricular blood that rapid fluctuation of the rate of change of pressure would appear in the carotid. High frequency distortions are often superimposed on undamped ventricular pressure records and perhaps too often dismissed as artefacts. But we found no evidence of HF distortion in the contour of the ballistocardiograms of any of these cases. The common types of cardiac incoordination produce distortions of these records of a lower frequency than those encountered in the carotid HF abnormality.

While other possibilities cannot be ruled out at this time the genesis of disturbed

flow in the carotid artery is clearly related to a lesion of the aortic valve in the great majority of our cases.

Conclusions

Records of the carotid pulse and its first time derivative have been taken, under standard resting conditions, in 349 cardiac patients, 180 of whom were subjected to cardiac catheterization. These results have been analyzed to throw light on an abnormality of pulse contour, conspicuous in the pulse derivative—a burst of high frequency distortion occurring at and after the peak of the main wave.

The presence of this abnormality is closely related to the presence of aortic valve lesions diagnosed either by the aortic valve gradient or by conventional clinical evidence. While such a lesion must be a prime factor in its genesis in most cases, the evidence also indicates that a second factor must be present before the abnormality is produced: sufficient velocity of the ejected blood. This theory is satisfactory in 98 per cent of our cases; the few discrepancies are discussed.

Such rapid fluctuations in the carotid pulse derivative indicate a disturbance of blood flow in which its normal streamlined character is broken up by vortices, jets, and eddies. Such a disturbance should not be called turbulence because turbulence is defined today as a phenomenon of much higher frequency than our methods would detect. The eddies and whorls of such disturbed flows must increase the impedance to cardiac ejection and like turbulence make the heart's task more difficult. This handicap is very common in aortic valve disease.

REFERENCES

- 1 Smith J E. A technique for recording carotid artery pulsations with special reference to aortic stenosis. *AM HEART J* 49:428 1955.
- 2 Conn H L Jr and Horwitz O. *Cardiac and vascular diseases*. Philadelphia 1971. Lea & Febiger Publishers.
- 3 Starr I and Ogawa S. On taking the first derivative of the pulse and its relations to the ballistocardiogram. *Fed Proc* 19:106 1960.
- 4 Starr I and Ogawa S. A clinical study of the first derivative of the brachial pulse. Normal standards and abnormalities encountered in heart disease. *AM HEART J* 65:482 1963.
- 5 Mason D T, Braunwald E, Ross J Jr and Morrow A G. Diagnostic value of first and second derivatives of the arterial pressure pulse in aortic valve disease and hypertrophic sub-aortic stenosis. *Circulation* 30:90 1964.
- 6 Boiteau G M and Allenstein B J. Hypertrophic sub-aortic stenosis. *Am J Cardiol* 8:614 1961.
- 7 Braunwald E, Goldblatt A, Aygen M M, Rockoff S D and Morrow A G. Congenital aortic stenosis: clinical and hemodynamic findings in 100 patients. *Circulation* 27:426 1963.
- 8 Brecht K and Bouche H L. Zur Abnahme des Arterienpulses am Menschen mit den Infrazton Mikrophon. *Pflügers Arch* 257:490 1953.
- 9 Ettinger E O. Flow patterns and vascular geometry. In Ettinger E O, editor. *Pulsatile blood flow*. New York 1964. McGraw Hill Book Company, Inc.
- 10 Side C D and Gosling R G. Non-surgical assessment of cardiac function. *Nature* 232:335 1971.
- 11 Schultz R D, Hokanson D E and Strandness D E Jr. Pressure flow and stress strain measurements of normal and diseased aortic segments. *Surg Gynecol Obstet* 124:1767 1967.
- 12 Mercer E N and Walters M B. Mitral regurgitation simulating aortic stenosis. *Can Med Assoc J* 93:413 1965.

Reciprocal movement of the right and left heart, demonstrated by directional Doppler ultrasound

Dennis Abelson MD

Hans R Muller MD

Philadelphia Pa and Basel Switzerland

The Doppler signals which can be detected over the precordium using an ultrasonic source have previously been described.^{1,2} If the difference frequency which lies in the audible range is presented irrespective of sign to a zero crossing detector a characteristic pattern of unidirectional spikes is produced corresponding to wall movement, valve leaflet movement and blood flow.³ The advent of directional equipment which has been used by several investigators⁴⁻¹⁰ to study peripheral blood flow non-invasively provided us with an opportunity to compare the movements of the right and left sides of the heart.

In the directional instrument⁴ Doppler shifted frequencies above the 4 MHz center frequency (ie movements towards the probe) are fed to one zero crossing detector while frequencies below 4 MHz (movements away from the probe) are fed to a second detector. The output of the latter is inverted after which the two are combined to produce a single differential output.

Observations were made on six normal

subjects in the supine position. One probe was placed on the precordium just internal to the apex pointing directly posteriorly while a second probe was located over the epigastrium just to the right of the xiphoid process and pointing upwards backwards and to the right—ie normal to the inferior surface of the right ventricle. Recordings were made simultaneously with the electrocardiogram on a Mingograph electroencephalograph.

During systole movement towards the probe was detected over the left ventricle. At the same time over the right ventricle reciprocal movement away from the probe was recorded. In diastole the direction of the movements was reversed. Fig 1 shows two representative tracings.

The deflections obtained over the apical area both in systole and diastole correspond in time and direction with the movement of the anterior surface of the heart. On the right side the probe is closer to the base of the heart which is tethered by its attachments to the great vessels. The inferior surface of the right ventricle would thus be expected to retreat from the anterior chest

From the Department of Research, The Lankenau Hospital, Philadelphia, Pa. and the Laboratory of ECG and Diagnostic Ultrasonography, University of Basel, Switzerland.
Received for publication: June 21, 1973.
Reprint request to Dennis Abelson MD, The Lankenau Hospital, 100 Locust Street, Philadelphia, Pa. 19105.
Ditmer Delandale Electrocardiography 92C, University of Basel.

flow in the carotid artery is clearly related to a lesion of the aortic valve in the great majority of our cases

Conclusions

Records of the carotid pulse and its first time derivative have been taken under standard resting conditions, in 349 cardiac patients 180 of whom were subjected to cardiac catheterization. These results have been analyzed to throw light on an abnormality of pulse contour, conspicuous in the pulse derivative, a burst of high frequency distortion occurring at and after the peak of the main wave.

The presence of this abnormality is closely related to the presence of aortic valve lesions, diagnosed either by the aortic valve gradient, or by conventional clinical evidence. While such a lesion must be a prime factor in its genesis in most cases the evidence also indicates that a second factor must be present before the abnormality is produced sufficient velocity of the ejected blood. This theory is satisfactory in 98 per cent of our cases the few discrepancies are discussed.

Such rapid fluctuations in the carotid pulse derivative indicate a disturbance of blood flow in which its normal streamlined character is broken up by vortices jets and eddies. Such a disturbance should not be called turbulence because turbulence is defined today as a phenomenon of much higher frequency than our methods would detect. The eddies and whorls of such disturbed flows must increase the impedance to cardiac ejection and like turbulence make the heart's task more difficult. This handicap is very common in aortic valve disease.

REFERENCES

- 1 Smith J E A technique for recording carotid artery pulsations with special reference to aortic stenosis *Am Heart J* 49:428 1955
- 2 Conn H L Jr and Horwitz O Cardiac and vascular diseases Philadelphia 1971 Lea & Febiger Publishers
- 3 Starr I and Ogawa S On taking the first derivative of the pulse and its relations to the ballistocardiogram *Fed Proc* 19:106 1960
- 4 Starr I and Ogawa S A clinical study of the first derivative of the brachial pulse. Normal standards and abnormalities encountered in heart disease *Am Heart J* 65:482 1963
- 5 Mason D T Braunwald E Ross J Jr and Morrow A G Diagnostic value of first and second derivatives of the arterial pressure pulse in aortic valve disease and hypertrophic sub-aortic stenosis *Circulation* 30:90 1964
- 6 Botteru G M and Allenstein B J Hypertrophic sub-aortic stenosis *Am J Cardiol* 8:614 1961
- 7 Braunwald E Goldblatt A Aygen M M Rockoff S D and Morrow A G Congenital aortic stenosis clinical and hemodynamic findings in 100 patients *Circulation* 27:426 1963
- 8 Brecht K and Bouche H L Zur Abnahme des Arterienpulses am Menschen mit den Infra- und Mikrophon *Pfluegers Arch* 261:490 1953
- 9 Ettinger E O Flow patterns and vascular geometry in Ettinger E O editor *Pulsatile blood flow* New York 1964 McGraw Hill Book Company Inc
- 10 Side C D and Gosling R G Non surgical assessment of cardiac function *Nature* 232:335 1971
- 11 Schultz R D Hokanson D E and Strandness D E Jr Pressure-flow and stress-strain measurements of normal and diseased aortic segments *Surg Gynecol Obstet* 124:1267 1967
- 12 Mercer E N and Walters M B Mitral regurgitation simulating aortic stenosis *Can Med Assoc J* 93:413 1965

The effect of noradrenaline on blood flow and oxygen consumption in normal and ischemic areas of myocardium

R J Marshall B Sc

J R Parratt B Pharm M Sc Ph D

Glasgow Scotland

Noradrenaline has been extensively used in the management of clinical shock of cardiac origin^{1,2} The most likely explanation for any clinical benefit achieved is that by elevating perfusion pressure noradrenaline increases collateral flow through a coronary vascular bed which is often atheromatous^{3,4} This would clearly be beneficial to patients whose myocardial performance is limited by a low coronary perfusion pressure Recently Maroko and associates⁵ have shown that elevating the systemic arterial pressure (with methoxamine) in dogs subjected to acute coronary artery occlusion markedly reduced ST segment elevation They concluded that elevating arterial pressure decreased the extent of ischemic injury presumably by elevating collateral flow To our knowledge however there have been no studies on the effects of noradrenaline on nutritive blood flow or on oxygen consumption in ischemic areas of the myocardium In the present study a technique has been used which enables simultaneous comparisons to be made of blood flow in ischemic and essentially normal areas of the left ventricular

wall Sampling blood from the coronary sinus (draining essentially normal areas of myocardium) and from a coronary vein (draining the ischemic muscle mass) also allowed simultaneous assessments to be made of oxygen consumption by these two areas

Methods

Nine greyhounds of both sexes and weighing between 25 and 34 kilograms were anesthetized with sodium thiopental (20 mg per kilogram of body weight injected intravenously) After endotracheal intubation anesthesia was maintained with 0.5 to 1.0 per cent trichloroethylene (Trilene) vaporized from a Tritec vaporizer (Cypreane Ltd) The dogs were on intermittent positive pressure ventilation with oxygen throughout the experiment the stroke volume of the Palmer ventilation pump was adjusted to give an expired end tidal PCO_2 between 26 and 38 mm Hg (arterial PCO_2 30 to 40 mm Hg 1 mm Hg = 1.333 mm bar) Reflex movement was prevented by the intermittent administration of succinylcholine (100 mg by intramuscular injection)

Received for publication January 30, 1973
Revised manuscript accepted for publication May 1, 1973
From the Department of Physiology, University of Glasgow, and the Department of Pharmacology, University of Glasgow, Glasgow, Scotland.
This work was supported by a grant from the Scottish Hospital Endowment Research Trust.
Reprint requests to Dr. J. R. Parratt, Department of Pharmacology, University of Strathclyde, Glasgow G1 1XH, Scotland.

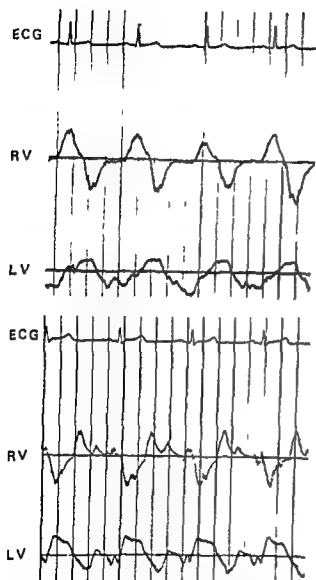


Fig. 1 top and bottom Simultaneous recordings of the electrocardiogram and directional velocity doppler cardiogram recorded over the right ventricle (RV) and left ventricle (LV) in two normal subjects. Movement toward the probe is shown as an upward deflection.

wall during systole, and approach it during diastole which is also in accord with the Doppler findings. Application of this non-invasive directional technique to disorders of cardiac movement is being pursued.

REFERENCES

- 1 Satomura S Ultrasonic Doppler method for the inspection of cardiac functions *J Acoust Soc Am* 29:1181 1957
- 2 Abelson D Ultrasonic Doppler auscultation of the heart with observations on atrial flutter and fibrillation *JAMA* 204:438 1968
- 3 Abelson D M Jaffe J and Murray P M Determination of cardiovascular velocities by dopplercardiometry I The normal tracing and effects of age *Cardiovasc Res* 5:535 1971
- 4 Strandness D E Jr Kennedy J W Judge T P and McLeod F D Transcutaneous directional flow detection A preliminary report *Am Heart J* 78 65 1969
- 5 Kalmanson D Veyrat C and Chiche P Venous return disturbances induced by arrhythmias *Cardiovasc Res* 4:279 1970
- 6 Folse R and Alexander R H Directional flow detection for localizing venous valvular incompetency *Surgery* 67:114 1970
- 7 Alexander R H Nippa J H and Folse R Directional transcutaneous assessment of venous inflow *Am Heart J* 82 86 1971
- 8 Reagan T R Miller C W and Strandness D E Jr Transcutaneous measurement of femoral artery flow *J Surg Res* 11 477 1971
- 9 Rittenhouse F A and Strandness D E Jr Oscillatory flow patterns in patients with aortic valve disease *Am J Cardiol* 28:568 1971
- 10 Muller H R The diagnosis of internal carotid artery occlusion by directional Doppler sonography of the ophthalmic artery *Neurology* 22 816 1972

Table 1 Hemodynamic effects of coronary artery ligation and of nor-adrenaline infusions (10 µg/kg)/min (mean ± S.E. n = 12)

| | Pre-ligation | 2 to 3 hr post ligation | Noradrenaline (2 to 3 hr post ligation) |
|--|--------------|-------------------------|---|
| Systolic blood pressure (mm Hg) | 158 ± 8 | 151 ± 9 | 218 ± 9† |
| Diastolic blood pressure (mm Hg) | 120 ± 5 | 111 ± 5 | 157 ± 6† |
| Mean blood pressure (mm Hg) | 135 ± 6 | 125 ± 7 | 177 ± 7 |
| Heart rate (beats/min) | 166 ± 5 | 188 ± 10 | 236 ± 21 |
| Cardiac output (L/min) | 3.2 ± 0.4 | 1.9 ± 0.2 | 2.4 ± 0.2 |
| Cardiac work (kg M/min) | 5.8 ± 0.8 | 3.4 ± 0.5 | 6.1 ± 0.7† |
| Peripheral vascular resistance (arbitrary units) | 49 ± 8 | 71 ± 8 | 85 ± 11 |
| LVEDP (mm Hg) | 10.1 ± 1.6 | 16 ± 3 | 20 ± 3 |
| Lv dP/dt max (mm Hg/s) | 1572 ± 154 | 1591 ± 157 | 3316 ± 277† |
| Left circumflex coronary flow (ml/min) | 90 ± 8 | 19 ± 8 | 143 ± 13† |
| Myocardial oxygen consumption (ml/min) | 11.7 ± 0.6 | 11.1 ± 1.3 | 20.1 ± 1.8† |
| Myocardial vascular resistance (arbitrary units) | 1.4 ± 0.2 | 1.6 ± 0.1 | 1.2 ± 0.1 |
| Total body oxygen consumption (ml/min) | — | 126 ± 10 | 112 ± 17 |

P < 0.05

† P < 0.0005

essentially normal region ml/min)
left circumflex flow (ml/min) × arterial — coronary sinus oxygen contents (ml/ml blood)

- (2) Myocardial oxygen consumption (ischemic region ml/100 Gm/min)
infarct flow (ml/min) × arterial — coronary vein oxygen content (ml/ml blood)
- (3) Myocardial vascular resistance (essentially normal myocardium arbitrary units)

diastolic blood pressure (mm Hg) — right atrial pressure (mm Hg)

mean left circumflex flow (ml/min)

- (4) Myocardial vascular resistance in the ischemic region was separated into two components—collateral (large vessel) network resistance and the resistance of the small vessels in the ischemic region itself (infarct resistance). They were calculated as follows

A collateral resistance (arbitrary units) diastolic blood pressure (mm Hg) — diastolic PCP (mm Hg)/PCF (ml/min)

B infarct resistance (arbitrary units) diastolic PCP (mm Hg) — right

atrial pressure (mm Hg)/ml xenon clearance (ml/100 Gm/min)

- (5) Whole body oxygen consumption (ml/min) cardiac output (ml/min) × arterial — right atrial oxygen content (ml/ml)

External cardiac work peripheral vascular resistance myocardial oxygen availability and oxygen extraction coefficients were calculated as previously described¹⁴ (*) Noradrenaline (10 µg per kilogram of body weight per minute as base was infused into a brachial vein using a LKB slow infusion pump (rate 1.0 ml/min) for a period of ten to fifteen minutes two to three hours after acute coronary artery ligation (mean time after ligation 2 hrs 35 min). Twelve experiments were performed in nine dogs. All values quoted in the text are mean ± S.E. Statistical analysis was performed using Student's paired t test.

Results

The effects of coronary artery ligation. Two to three hours after acute coronary artery ligation the most pronounced hemodynamic changes were decreases in cardiac output and in external cardiac work (Table I). Systemic arterial pressure was maintained despite the reduction in cardiac output by a compensatory increase in peripheral vascular resistance. Although left ventricular dP/dt max was unchanged the

Temperature was measured in the mid esophagus using a direct recording thermocouple (Ellab, Copenhagen)

Using a Siemens image intensifier, catheters were positioned under fluoroscopic control in the descending aorta, left ventricle, right atrium and coronary sinus and were used for blood sampling and for pressure measurements using Elema Schonander capacitance transducers. The rate of change of left ventricular pressure (dP/dt) was measured using an Elema Schonander differentiator circuit and for the measurement of left ventricular end diastolic pressure (LVEDP) the left ventricular pressure pulse was also recorded at high gain and the record cut off above 40 mm Hg. Cardiac output was measured by dye dilution. Indocyanine green (2.5 mg) was injected into the right atrium and blood was withdrawn from the descending aorta through a densitometer (Waters Co. Rochester, Minn.)

The heart was exposed through a left thoracotomy the pericardium was incised and the anterior descending branch of the left coronary artery was prepared for ligation at a position distal to the main septal branch. The anterior coronary vein (or one of its major branches) was catheterized using a Longdwell 4 inch Teflon catheter (20G). The catheter was manipulated so that the tip lay well into the potentially ischemic area; it was not found necessary to tie this catheter in position. After coronary artery ligation this catheter has been shown to drain blood predominantly from the infarcting area.^{7,8} Anaerobic blood samples (2 ml) were simultaneously obtained from this catheter and from the coronary sinus, right atrium and descending aorta and were analyzed for PO_2 , P_{CO_2} and pH using appropriately calibrated electrode systems (Radiometer, Copenhagen) as outlined by Ledingham and associates.⁹ To allow for any difference in the measurement of oxygen tension between gas and blood a blood gas factor was derived for each experiment using blood tonometered with a known tension of oxygen. Oxygen and carbon dioxide tensions and pH were corrected for any difference in temperature between the electrode and the animals mid esophageal temperature using the Radiometer blood gas calculator¹⁰ or an ap-

propriately programmed Hewlett Packard desk computer. Hemoglobin was measured using a co-oximeter (Instrumentation Laboratory Inc., Lexington, Mass.) and blood oxygen content was calculated using the formula

$$Hb (Gm) \times 1.39 \times \% \text{ saturation} / 100 + PO_2 \\ (\text{mm Hg}) \times 0.0031$$

There is a good correlation between blood oxygen content calculated by this indirect method and by simultaneous Van Slyke determinations.⁹

Blood flow in the left circumflex coronary artery was measured using a Nycotron 372 electromagnetic flowmeter and an implantable 2 or 2.5 mm probe. The probe was calibrated at the end of the experiment by perfusing the vessel with the animals own blood at known flow rates.

After a one to two hour stabilization period during which time several samples were taken for blood gas determinations from each of the four catheter sites (coronary sinus and vein, right atrium and aorta) an intravenous injection of lignocaine (20 to 40 mg) was given and the left anterior coronary artery was ligated in one stage. A Portex nylon catheter (O.D. 1.34 mm) was inserted distally into the artery and tied in position. This catheter was used for the assessment of peripheral coronary pressure (PCP) using a capacitance transducer for the measurement of back (retrograde or peripheral) coronary flow (PCF) from the open catheter and for the injection of small amounts (about 90 μ Ci, volume 1.2 ml) of ¹³³Xenon into the ischemic region. The clearance of radioactivity from the ischemic muscle was measured using a collimeter with a narrow angle of acceptance positioned directly over the ischemic region. This method of assessing nutritive blood flow in ischemic regions of the myocardium has been used by a number of workers including Rees and Redding¹¹ and Gregg¹² and Pasyk and associates.¹³

Left circumflex blood flow (phasic and mean) systemic arterial right atrial, left ventricular and peripheral coronary pressures left ventricular dP/dt and the electrocardiogram (standard limb Lead II) were recorded on an eight channel Elema Schonander ink jet writing recorder.

The following data were derived

(1) Myocardial oxygen consumption (cs

Table II Changes in the ischemic myocardium induced by noradrenaline
(mean \pm SE $n = 12$)

| | Control | Noradrenaline |
|---|----------------|-----------------|
| Peripheral coronary pressure (systolic mm Hg) | 53 \pm 4 | 71 \pm 7† |
| (diastolic mm Hg) | 18 \pm 2 | 33 \pm 4† |
| (mean mm Hg) | 35 \pm 3 | 56 \pm 4† |
| Peripheral coronary flow (ml/min) | 14 \pm 0.2 | 3.5 \pm 0.5† |
| Collateral resistance (arbitrary units) | 81 \pm 13 | 43 \pm 7 |
| Infarct blood flow (ml/100 Gm/min) | 17.6 \pm 2.7 | 39.6 \pm 4.3† |
| Infarct resistance (arbitrary units) | 2.2 \pm 0.3 | 1.6 \pm 0.2 |
| Infarct oxygen consumption (ml/100 Gm/min) | 2.8 \pm 0.5 | 5.3 \pm 0.8† |

P < 0.05

†P < 0.0005

administration of noradrenaline precipitated irregularities of cardiac rhythm. These consisted of nodal rhythms (6 experiments) or of bursts of ventricular ectopic activity (6 experiments). Reversion to sinus rhythm invariably occurred during the infusion period and in these animals it was possible to demonstrate that noradrenaline increased sinus activity and sometimes reduced ST segment depression. In all except one experiment left circumflex flow was markedly elevated during the period of the infusion and there was a substantial increase in normal myocardial consumption (Table I). Noradrenaline did not change coronary sinus PO_2 (29 ± 1 to 30 ± 2 , $P > 0.5$) or the oxygen extraction coefficient (32 ± 2 to 31 ± 3 per cent, $P > 0.9$). The ratio of oxygen available to the normal myocardium (flow \times arterial oxygen content) to oxygen consumed by the myocardium was unchanged.

The hemodynamic effects of noradrenaline are illustrated in Fig. 1.

The effect of noradrenaline on blood pH and gas tensions. A consistent and significant finding was a decrease in arterial pH during the noradrenaline infusion (7.439 ± 0.024 to 7.379 ± 0.023 units, $P < 0.001$). This was reflected in similar changes in coronary sinus and coronary vein pH (coronary sinus 7.376 ± 0.023 to 7.322 ± 0.024 units, $P < 0.001$; coronary vein 7.337 ± 0.025 to 7.313 ± 0.023 units, $P < 0.01$) and was almost entirely due to a small but significant elevation in arterial PCO_2 (32 ± 2 to 35 ± 2 mm Hg, $P < 0.01$). Arterial base deficit was not significantly altered by

noradrenaline (-5.8 ± 1.7 to -8.2 ± 1.2 mEq/L, $P > 0.01$). In our experience a base deficit of -5 mEq/L is normal in dogs anesthetized with trichloroethylene. Calculation of the pulmonary shunt ratio by the method described by Hyde¹⁸ indicated that noradrenaline increased shunting in this experimental preparation from a mean of 17 ± 2 per cent of the cardiac output to 24 ± 2 per cent. This change was significant ($P < 0.001$). However arterial PO_2 was not significantly altered by noradrenaline (272 ± 23 mm Hg before and 194 ± 40 during the infusion, $P > 0.1$). All the above changes were readily reversible within 10 to 20 minutes of stopping the infusion.

The effects of noradrenaline in the ischemic myocardium. Some of these results are summarized in Table II. Noradrenaline markedly increased peripheral coronary pressure and there was some evidence that this directly resulted from the elevated systemic arterial pressure (Fig. 2). Flow in the ischemic region as measured by xenon clearance (Fig. 3) or by retrograde flow (PCF) was markedly increased in all except one of the twelve experiments. The exception was the same animal in which there was no change in left circumflex flow in response to noradrenaline and it may be significant that this dog had an abnormally acid arterial pH of 7.290 units and base deficit of -16 mEq/L. There were substantial increases in oxygen consumption by the ischemic myocardium during noradrenaline administration (Table II) but no change in either oxygen extraction

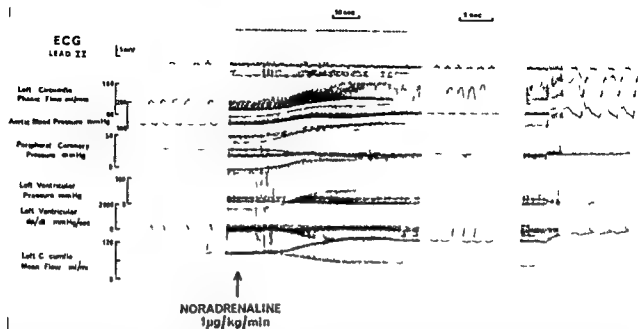


Fig. 1 The effects of an infusion of noradrenaline ($1 \mu\text{g/kg/min}$) commencing at the arrow on (from above) the electrocardiogram (Lead II) left circumflex coronary blood flow (phasic ml/min) aortic pressure (mm Hg) peripheral coronary pressure (mm Hg) left ventricular pressure (mm Hg) and dP/dt (mm Hg/sec) mean circumflex flow (ml/min) in a dog whose left anterior descending coronary artery had been ligated 24 hours previously. The right hand portion of the trace was recorded towards the end of the 12 min infusion period.

left ventricular end diastolic pressure (LVEDP) was significantly elevated (from 10.1 ± 1.6 to 16 ± 3 mm Hg) this unchanged LV dP/dt with an increased preload indicates a reduction in myocardial contractility. This is presumably because of the functional loss of a considerable portion (25 to 30 per cent) of the left ventricular wall.¹³ The area supplied by the ligated vessel was cyanosed, bulging and clearly non contractile and there was electrocardiographic evidence (ST segment depression) of myocardial ischemia. Flow in this ischemic region was 17.6 ± 2.7 ml/100 Gm/min, which corresponds to a reduction of about 80 per cent of the flow in this area prior to ligation.⁷ In contrast there was no change in blood flow to the normal region of the left ventricular wall or in oxygen consumption by the normal myocardium. The hemodynamic effects of acute coronary artery ligation are summarized in Table I.

The marked changes in local venous blood draining an acutely ischemic region in this experimental model have been outlined by Fisher and colleagues.⁷ There is a substantial production of lactate, a marked increase in PCO_2 and decreases in pH and

oxygen content. These changes are however transient and at the time that noradrenaline was administered (2 to 3 hr post ligation) there were again no differences in PO_2 , PCO_2 or pH between the coronary sinus blood (draining essentially normal areas of the myocardium) and coronary venous blood (draining the ischemic area). Before ligation the values for coronary venous blood were PO_2 31 ± 2 mm Hg, PCO_2 44 ± 2 mm Hg, pH 7.353 ± 0.014 units and the oxygen extraction coefficient 52 ± 3 per cent. The corresponding values 2 to 3 hr after ligation were PO_2 27 ± 1 mm Hg, PCO_2 47 ± 3 mm Hg, pH 7.337 ± 0.025 and 59 ± 3 per cent. Comparable values for coronary sinus blood were PO_2 32 ± 2 mm Hg, PCO_2 44 ± 2 , pH 7.364 ± 0.014 units and the oxygen extraction coefficient 50 ± 3 per cent and 2 to 3 hr after ligation 29 ± 1 mm Hg, PCO_2 44 ± 3 mm Hg, pH 7.356 ± 0.023 and 52 ± 2 per cent.

Hemodynamic effects of noradrenaline. When administered 2 to 3 hr after acute coronary artery ligation noradrenaline ($1.0 \mu\text{g}$ per kilogram of body weight per minute) markedly elevated systemic arterial pressure, cardiac work and LV dP/dt (Table I). In each of the twelve experiments the

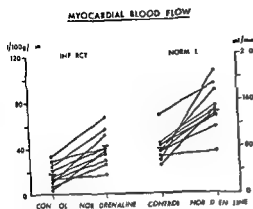


Fig 3 The effect of noradrenaline ($10 \mu\text{g/kg/min}$) on blood flow (ml/min) in normal myocardium and in a region supplied by a ligated coronary artery (infarct blood flow $\text{ml}/100 \text{ Gm}/\text{min}$)

preparations where PCP was only slightly elevated. This would seem to indicate an active vasodilator mechanism in at least some of the experiments. There is however no need to postulate a direct vasodilator action of noradrenaline in the ischemic myocardium. Vasodilation is more likely to be due to the elevated perfusion pressure washing out metabolites from the ischemic muscle and this could account for the decreased vascular resistance in the infarct. Noradrenaline certainly increased the carbon dioxide tension (from 47 ± 3 to 53 ± 3 $P < 0.02$) and decreased the pH (from 7.337 ± 0.02 to 7.313 ± 0.023 units $P < 0.05$) of blood draining the ischemic area. We believe however that such active vasodilator changes are of minor importance. The major factor increasing flow in the ischemic area is the elevated perfusion pressure. This would be in agreement with the finding of Maroko and colleagues⁶ that the extent of ischemia is highly dependent upon systemic blood pressure. Elevating systemic pressure (with methoxamine) decreased the extent of ischemic muscle injury while decreasing systemic pressure (by hemorrhage) increased both the area and the severity of ischemia.

It has not been possible in this study to separate flow changes in the epicardial and endocardial regions of the ischemic area. Partial coronary occlusion is accompanied by a disproportionate decrease in endo-

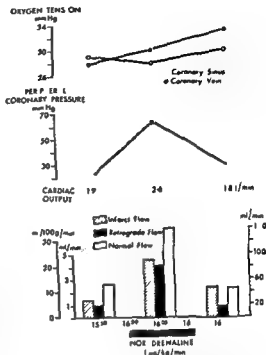


Fig 4 The effect of noradrenaline ($10 \mu\text{g/kg/min}$) in a dog 2 to 3 hr after coronary artery ligation. From above: coronary sinus and vein Po_2 (mm Hg), mean peripheral coronary pressure (mm Hg), normal myocardial blood flow (\square ml/min), retrograde flow from the ligated vessel (\blacksquare ml/min) and flow within the ischemic region (infarct flow \blacksquare $\text{ml}/100 \text{ Gm}/\text{min}$). The artery was ligated at 13:30 hr. Noradrenaline (administered at time 16:30 hr to 18:00 hr) markedly elevated flow both in the ischemic region and in the essentially normal myocardium.

cardial flow^{18,20} and this is probably the explanation for the more extensive necrosis which occurs in subendocardial layers after coronary artery occlusion.²¹ Postmortem sectioning of the ischemic region showed clearly that this was also the case in the present experiments. Fulton²² has postulated that blood flow into the subendocardial plexus (supplying the inner layers of the myocardium) only occurs during diastole. This means that the effective driving pressure for the endocardial region is the difference between the pressure in the coronary artery during diastole and the end diastolic pressure in the left ventricle. This would normally be about 110 mm Hg (Table I). After coronary ligation however this is drastically reduced since it then becomes the difference between diastolic

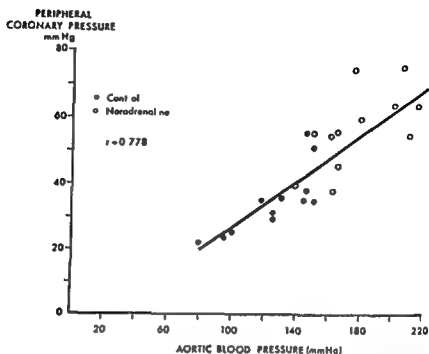


Fig 2 The relationship between systemic aortic blood pressure (mm Hg) and peripheral coronary pressure (mm Hg) before (●) and after (○) noradrenaline in dogs subjected to acute coronary artery ligation

(59 ± 3 per cent before noradrenaline and 51 ± 4 per cent during the infusion) or in coronary vein PO_2 (27 ± 1 mm Hg to 31 ± 3 mm Hg $P > 0.05$). The ratio of oxygen available to the ischemic area to oxygen consumed by the ischemic area was unchanged by noradrenaline.

Some attempt was made to calculate vascular resistance changes both in the normal myocardium and in the ischemic region. Vascular resistance in the normal myocardium was slightly ($P < 0.02$) reduced by noradrenaline. This was so whether resistance was calculated from mean flow or from flow during diastole. Resistance in the ischemic region was separated into 'collateral network resistance' (the resistance to flow of the collateral vessels supplying the ischemic region) and 'infarct resistance' (the resistance of the smaller vessels of the ischemic region itself). Noradrenaline decreased resistance to flow both in the infarct and, more markedly in the collateral network (Table II) and increased the pressure gradient across the collateral network (mean systemic pressure—mean PCP) from 90 to 121 mm Hg.¹² Some of the effects of noradrenaline on the ischemic myocardium are illustrated in Fig 4.

Discussion

These results demonstrate that noradrenaline increases blood flow through a region of myocardium made ischemic by the ligation of a major branch of the left coronary artery. Such an increase in flow could be due to the increased perfusion pressure (in this case the PCP) and/or to an active drug induced vasodilator mechanism. In the normal myocardium it is generally agreed that with noradrenaline both mechanisms operate.¹⁷ There is both an increased perfusion pressure and also active vasodilation. The latter is secondary to the raised myocardial oxygen consumption which results from increased contractility. It is likely that both mechanisms also operate in the ischemic myocardium. If the increase in nutritional blood flow (125Xenon clearance) resulted solely from an elevated perfusion pressure one would expect a linear relationship between these variables. In general when PCP is plotted against infarct blood flow (Fig 5) there is a good relationship. There is however a greater scatter when the noradrenaline results are plotted on the same graph and, although over all the flow/perfusion pressure relationship was still linear, considerable increases in 125Xenon clearance occurred in

external cardiac work and an increase in left ventricular end-diastolic pressure with an unchanged LV dP/dt. This is indicative of reduced myocardial contractility. Blood flow in the area supplied by the ligated vessel fell to a mean of 17.6 ± 2.7 ml/100 Gm/min which is about 20 per cent of the normal flow in this region.

Infusing noradrenaline 2 to 3 hr after ligation increased systemic arterial pressure LV dP/dt max external cardiac work and blood flow and oxygen consumption in normal areas of the myocardium. There was some evidence that pulmonary shunt flow was increased by the drug.

Noradrenaline also markedly increased peripheral coronary pressure and blood flow in the ischemic region as assessed by ^{133}Xe clearance and by retrograde flow from the ligated vessel. Oxygen consumption in the ischemic region was also increased by noradrenaline. It is suggested that part of the increase in flow occurs in the endocardial region since the effective subendocardial perfusion pressure is increased by noradrenaline. This increase in flow would account for the reduction in infarct size which has been observed by other workers when the systemic arterial pressure is raised.

REFERENCES

- 1 Nies A S and Melmon K L The rational management of cardiogenic shock in Brest A N ed Cardiovascular therapy Philadelphia 1969 F A Davis Company pp 65-88
- 2 Perlothe M G and Harrison D C Cardiogenic shock: a review Clin Pharmacol Ther 10:499 1969
- 3 Corday E Bazika V and Lann T W Vasopressor treatment of cardiogenic shock in Mills L C and Moyer J H editors Shock and hypotension Pathogenesis and treatment New York 1965 Grune & Stratton Inc pp 526-536
- 4 Mueller H Ayres S M Gregory J J Giannelli S and Grace W J Hemodynamics coronary blood flow and myocardial metabolism in coronary shock: response to 1 norepinephrine and isoproterenol J Clin Invest 49:1885 1970
- 5 Mueller H Ayres S M Giannelli S Conklin E F Mazzara J T and Grace W J Effect of isoproterenol 1 norepinephrine and intraaortic counterpulsation on hemodynamics and myocardial metabolism in shock following acute myocardial infarction Circulation 44:335 1971
- 6 Marolo I R Kjekshus J K Sobel B E Watanabe T Covell J W Ross J and Braunwald E Factors influencing infarct size following experimental coronary artery occlusions Circulation 43:67 1971
- 7 Fisher W Heimbach D Ledingham J McAl Marshall R J and Parratt J R Blood flow and oxygen extraction in ischaemic and normal regions of the myocardium following acute coronary artery ligation J Physiol (Lond) 240:15P 1973
- 8 Marshall R J Parratt J R and Ledingham J McAl Unpublished data
- 9 Ledingham J McAl McBride T J Parratt J R and Vance J I The effect of hypercapnia on myocardial blood flow and metabolism J Physiol (Lond) 210:87 1970
- 10 Severinghaus J W Blood gas calculator J Appl Physiol 21:1108 1966
- 11 Rees J H and Redding V J Anastomotic blood flow in experimental myocardial infarction: A new method using ^{133}Xe clearance for repeated measurements during recovery Cardiovasc Res 1:169 1967
- 12 Gregg D E The microcirculation of the heart in reduced flow states in Shepro D and Fulton C I editors Conference on the microcirculation as related to shock Boston University 1967 London 1968 Academic Press pp 51-67
- 13 Pasik S Bloor C M Akhoury M and Gregg D E Systemic and coronary effects of coronary artery occlusion in the unanesthetized dog Am J Physiol 220:646 1971
- 14 Ledingham J McAl Parratt J R Smith G and Vance J P Haemodynamic and myocardial effects of hyperbaric oxygen in dogs subjected to haemorrhage Cardiovasc Res 5:277 1971
- 15 Marshall R J and Parratt J R Unpublished data
- 16 Hyde R W Clinical interpretation of arterial oxygen measurements Med Clin North Am 54:617 1970
- 17 Parratt J R Pharmacological aspects of the coronary circulation in Ellis G P and West G B editors Progress in medicinal chemistry London 1969 Butterworth & Co Ltd 6:111-66
- 18 Mour T W and DeBra D W Effect of left ventricular hypertension ischemia and vasoactive drugs on the myocardial distribution of coronary flow Circ Res 21:65 1967
- 19 Gregg D M and Nakamura Y Effect of coronary constriction on myocardial distribution of iodoantipyrine ^{133}I Am J Physiol 215:1082 1968
- 20 Becker L C Fortuin N J and Litt B Effect of ischemia and antianginal drugs on the distribution of radioactive microspheres in the canine left ventricle Circ Res 28:263 1971
- 21 Jennings J B Sommers H M Smyth G A Flack H A and Linn H Myocardial necrosis induced by temporary occlusion of a coronary artery in the dog Arch Pathol 70:68 1960
- 22 Fulton W F M The coronary arteries Springfield Ill 1965 Charles C Thomas Publisher

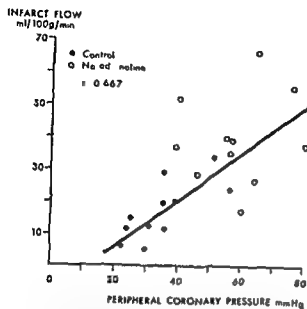


Fig 5 The relationship between infarct blood flow (125 I xenon clearance ml/100 Gm/min) and peripheral coronary pressure (mm Hg) in nine dogs 2 to 3 hr after coronary artery ligation before (●) and during (○) the infusion of noradrenaline

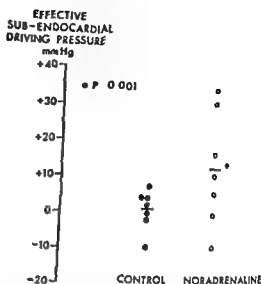


Fig 6 The effect of noradrenaline ($10 \mu\text{g/kg/min}$) on the effective sub endocardial driving pressure (mm Hg calculated a diastolic PCP minus LVEDP see discussion) The control values are those for the ischemic region obtained immediately prior to the noradrenaline infusions

peripheral coronary pressure and LVEDP. The left ventricular end diastolic pressure, as we have seen (Table I) elevated after ligation. Such calculated effective sub endocardial driving pressures are shown in Fig 6. It can be concluded from these experiments that there is hardly any pressure head for flow in the inner (deep) muscle layers of the developing infarct. Noradrenaline, however, substantially increased the effective driving pressure (Fig 6). It is likely therefore that part of the increased flow, which occurs through the total ischemic area during the administration of noradrenaline represents flow through the subendocardium. Thus although there is no evidence that noradrenaline increases the ratio of LV endocardial flow to LV epicardial flow in the normal myocardium,¹⁸ these present results might suggest that like propranolol²⁰ the ratio of LV endocardial/LV epicardial flow is increased by noradrenaline in the ischemic myocardium. An alternative way of increasing LV endocardial perfusion would be to reduce LVEDP. This has been suggested as the possible mechanism through which the β adrenoceptor drug oxyfedrine increases flow in developing infarcts.^{22,24}

The finding that noradrenaline increases tissue perfusion in ischemic muscle would explain the finding, in patients in shock

following acute myocardial infarction that myocardial lactate production is shifted to extraction during noradrenaline administration.⁵ It would also explain the finding that elevating systemic pressure is a factor which leads to a reduction in infarct size following experimental coronary artery occlusion in dogs.⁶ The fact that infarct oxygen consumption is increased by noradrenaline in our experiments suggests that within the infarcting myocardium there are viable as well as dead cells and that noradrenaline by restoring oxygen availability to these potentially viable cells reduces cell death and decreases the area of necrosis.

Summary

The effects of infusions of noradrenaline ($10 \mu\text{g/Kg/min}$) were studied in dogs 2 to 3 hr after acute ligation of the anterior descending branch of the left coronary artery. The model allowed blood flow to be simultaneously measured in the ischemic (infarcting) region and in the normal myocardium. Sampling blood from a local vein (draining the ischemic region) and from the coronary sinus (draining the normal myocardium) allowed comparisons to be made of oxygen consumption by the two regions.

Coronary artery ligation resulted in a marked decrease in carotid output and

The effect of heroin and multiple drug abuse on the electrocardiogram

Janet Lipski MD

Barry Stummel MD

Ephraim Donoso MD

New York N Y

Sudden death following parenteral injection of heroin is not a rare occurrence. Although the term narcotic overdose has been used to describe the sudden death phenomenon, it is now realized that in many instances the concentration of heroin in the tissues or blood is insufficient to cause toxicity and the precise mechanism of death is not clear.¹

It is not unreasonable that the sudden death syndrome seen in heroin users may be similar to those occurring in those with electrocardiographic abnormalities and are related to cardiac arrhythmias. The following study was undertaken in an attempt to define the effects of street heroin and other drugs of abuse on the electrocardiogram.

Materials and methods

Seventy-five asymptomatic individuals admitted to a methadone maintenance treatment program (VATP) served as the study group. All patients received a complete history and physical examination including a chest x-ray. A 12-channel SMA (sequential multiple analyzer) was performed as well as serum analysis for sodium, potassium, chloride, CO₂ content, urea nitrogen, and glucose. Urine analysis with

thin layer chromatography² was performed to detect the presence of morphine (heroin), methadone, quinine, barbiturates, and amphetamines. Routine and microscopic urine analysis was also performed. Any individual with cardiac, pulmonary, or renal disease by either history, physical examination, or laboratory findings was excluded from the study. A standard 12-lead ECG was obtained after a 20-minute rest period and was analyzed for abnormalities. All QT intervals were corrected for heart rate according to the tables of Ashman and Hull.³

Two groups of patients were studied. Group I (34 patients) had electrocardiograms while on parenteral heroin prior to initiation of methadone therapy. All gave a history of having used heroin confirmed by urine analysis within a 24-hour period of their ECG. Serum was drawn for the appropriate laboratory studies upon completion of the ECG. Group II comprised 41 patients who had been on methadone maintenance for varying periods of time but were still using other drugs sporadically, notably alcohol, barbiturates, heroin, or cocaine. All of these patients were found to have barbiturates in their urine or gave

From the Department of Medicine, Division of Cardiology, The Mount Sinai School of Medicine of The City University of New York, New York.
Received for publication February 5, 1973.
Reprint requests to Dr. Janet Lipski, Department of Medicine, Division of Cardiology, The Mount Sinai Hospital, 100th St., New York, N.Y. 10029.

- 23 Ledingham I McA McArdle C and Parratt J R Comparison of a coronary vasodilator drug (carbochromen) and a cardiac stimulant (oxyfedrine) on blood flow and oxygen extraction in experimental myocardial infarction Br J Pharmacol 44 323 1972
- 24 Parratt J R and Ledingham I McA Ex

perimental infarction in the dog changes in blood flow and in oxygen extraction induced by oxyfedrine in areas of developing myocardial ischaemia in Gerlach E and Moser K editors Wirkungsweise von Oxyfedrine Stuttgart and New York 1972 Schattauer Verlag pp 201 207

Table IV Correlation of heart rate QT interval and urine analysis (Group I patients)

| Patient no and initials | Heart rate | QT | QT | Urine analysis | | | | |
|-------------------------------|---------------|----|----|----------------|-----------|-------------|-------------|---------|
| | | | | Morphine | Methodone | Barbiturate | Amphetamine | Quinine |
| 1 C A | 102 | 40 | † | 0 | + | 0 | 0 | + |
| 2 C J | 70 | 42 | † | + | + | 0 | 0 | 0 |
| 3 H S | 75 | 38 | | + | + | 0 | 0 | + |
| 4 H D | 60 | 43 | | + | 0 | 0 | 0 | 0 |
| 5 I N | 65 | 38 | | + | 0 | 0 | 0 | 0 |
| 6 I J | 65 | 38 | | + | 0 | 0 | 0 | + |
| 7 J A | 104 | — | | + | 0 | 0 | 0 | 0 |
| 8 I S | 60 | 36 | | + | 0 | 0 | 0 | 0 |
| 9 M L | 75 | 37 | | + | 0 | 0 | 0 | + |
| 10 R F | 48 | 40 | | + | 0 | 0 | 0 | 0 |
| 11 R J | 85 | 38 | | + | 0 | 0 | 0 | 0 |
| 12 R H | 80 | 40 | | + | 0 | 0 | 0 | 0 |
| 13 S J | 100 | 32 | | + | 0 | 0 | 0 | + |
| 14 S P | 75 | 36 | | + | + | 0 | 0 | 0 |
| 15 S C | 110 | 30 | | + | + | 0 | 0 | 0 |
| 16 T O | 75 | 40 | † | + | + | 0 | 0 | 0 |
| 17 V A | 70 | 38 | | 0 | + | 0 | 0 | 0 |
| 18 V P | 98 | 32 | | 0 | 0 | 0 | 0 | 0 |
| 19 V D | 78 | 36 | | 0 | 0 | 0 | 0 | 0 |
| 20 S G | 71 | 36 | | + | 0 | 0 | 0 | 0 |
| 21 A M | 68 | 37 | | + | + | 0 | 0 | 0 |
| 22 B J | 62 | 39 | | + | 0 | 0 | 0 | 0 |
| 23 D L | 93 | 40 | † | + | 0 | 0 | 0 | 0 |
| 24 C N | 68 | 37 | | + | 0 | 0 | 0 | + |
| 25 M M | 70 | 38 | | + | + | 0 | 0 | + |
| 26 O P | 58 | 37 | | + | 0 | 0 | 0 | 0 |
| 27 P D | 52 | 42 | | 0 | + | 0 | 0 | 0 |
| 28 R R | 70 | 41 | † | 0 | + | 0 | 0 | 0 |
| 29 R D | 72 | 38 | | 0 | 0 | 0 | 0 | + |
| 30 R S | 83 | 36 | | + | 0 | 0 | 0 | 0 |
| 31 S V | 62 | 40 | | 0 | + | 0 | 0 | 0 |
| 32 S F | 58 | 36 | | + | + | 0 | 0 | + |
| 33 T R | 57 | 36 | | + | + | 0 | 0 | + |
| 34 W A | 64 | 42 | † | 0 | + | 0 | 0 | 0 |

† Corrected QT interval values for rate that are increased as per Ashman and *et al.*

postmortem were negative for all abusive drugs.¹

The precise mechanism of death is not clear. Those postulated include a pharmacologic overdose, an allergic type reaction to heroin, acute reactions to injected bacteria, quinine or other foreign material, anoxia, synergistic effects from multiple drug abuse, and pulmonary edema.^{5,7} Pulmonary edema in patients with acute intoxication has been recognized for nearly a hundred years,⁸ though its pathogenesis remains obscure.

The sudden onset and rapid fatal termi-

nation of the acute reaction with the needle found remaining in the vein in some instances¹ allows for the possibility of a fatal arrhythmia being the mode of exodus in these patients.

Heroin is three to four times more potent than morphine and when injected is hydrolyzed rapidly to monoacetyl morphine and subsequently to morphine. It is excreted in the urine largely as free and conjugated morphine. Morphine and related narcotics act by increasing vagal tone and by inhibiting the hydrolysis of acetylcholine.⁹ The effect of morphine on heart

Table I Age distribution of patients

| Age | Group I | Group II |
|-------|---------|----------|
| 10-19 | 0 | 0 |
| 20-29 | 18 | 15 |
| 30-39 | 13 | 19 |
| 40-49 | 3 | 6 |
| 50-59 | 0 | 1 |
| F/M* | 7/27 | 9/32 |

*Female/male ratio

Table II Cardiac abnormalities in patients in Group I

| Cardiac abnormality | No |
|-------------------------------------|----|
| Bradyarrhythmia | 6 |
| QT prolongation | 6 |
| Sinus tachycardia | 2 |
| Intra atrial conduction disturbance | 2 |
| Abnormal T waves | 2 |

Table III ECG changes in 15 patients taking heroin within 4 hours of examination (Group I)

| Abnormality | No |
|----------------------------|----|
| Tachycardia or bradycardia | 7 |
| Increased QT | 2 |

a history of drug abuse within a three day period of their electrocardiogram

The ECG's of 32 patients (Group III) free of cardiac pulmonary or renal disease matched for age and ethnic background were obtained from a routine employee health screening clinic. These individuals did not have a history of drug abuse.

Results

There were no serum or routine urine abnormalities in any group. The age distribution of patients is shown in Table I. The average age of Group I was 30 years, Group II, 33 years and Group III 34 years. The female/male ratio was approximately 1 to 4 in each group.

Of the 34 patients in Group I (only on heroin) abnormalities of the electrocardiogram were noted in 19 patients (55 per cent) (Table II). Of 15 patients who had taken heroin within 4 hours of their ECG changes were noted in 10 (68 per cent) (Table III). The most frequent abnormality in this group was a prolonged QT interval (19 per cent) and bradyarrhythmias (sinus arrhythmias with slow ventricular response or sinus bradycardia) (19 per cent). Correlation of heart rate, QT interval and urine analysis in Group I patients is seen in Table IV. Of the 41 patients on methadone with a history of multiple drug abuse, Group II, 27 (66 per cent) had some ECG changes. QTc prolongation was noted in 14 (34 per cent), prominent U waves in 13 (32 per cent) and bradyarrhythmias in 13 (32 per cent) (Table V). The most striking electrocardiogram in this group is from a 31 year-old addict of 17 years duration who died suddenly on the street. He was on 100 mg of methadone a day and had an impressive barbiturate habit as well as a history of sporadic parenteral cocaine use. The ECG (Fig. 1) taken several days prior to his death revealed marked prolongation of the QTc interval 0.48 sec with U waves of greater amplitude than the T waves. Postmortem examination by the medical examiner's office revealed a fresh puncture site but no gross abnormalities of the heart or lungs. Subsequent toxicological reports were negative except for the presence of methadone.

In the control group sinus bradycardia was present in 4 patients (12.5 per cent) and none had QT prolongation.

Discussion

Mortality rate related to heroin addiction has been increasing steadily. In New York City it was reported that narcotics chiefly heroin were the leading cause of death in 1969 and in 1970 in all males aged 15 to 35 including non addicts.⁴ An acute drug reaction was the cause of death in more than three quarters of all fatalities from narcotics; however a pharmacologic lethal level could not be demonstrated in most cases. In some toxicological studies at

Table V Cardiac abnormalities in patients in Group II

| Cardiac abnormality | No |
|--|----|
| QT prolongation | 14 |
| Prominent U wave | 13 |
| Bradyarrhythmias | 13 |
| Dimpled T wave | 7 |
| Abnormal T wave (Low voltage inversion peaked) | 5 |
| Abnormal LAD | 2 |
| First degree heart block | 2 |
| Intra atrial conduction abnormality | 2 |
| Premature systoles (apc a pc s) | 3 |
| Tachycardia | 1 |

In our series of patients transient atrial fibrillation was not found. However bradyarrhythmias and repolarization abnormalities were common in both groups (Table VI). Bradycardia enhances ventricular ectopic activity lowers the threshold to ventricular fibrillation^{21,22} and may serve as an important factor in precipitating ventricular tachycardia fibrillation and sudden death.

Prolongation of the QT interval represents delayed ventricular repolarization and an increase in the duration of the vulnerable period with greater susceptibility to the development of a ventricular dysrhythmia. In the presence of the QT prolongation single or multiple ventricular premature beats may lead to ventricular fibrillation and death. In addition it has been found that excitement and fright usually cause the QT interval to prolong further²³. In the addict usually operating under severe stressful situations additional lengthening of an already prolonged QT interval may well be responsible for ventricular fibrillation and death.

In these individuals with multiple drug abuse the hypotension produced by barbiturates,² respiratory depression produced by methadone²⁴ and the increased incidence of ventricular arrhythmias associated with alcohol²⁵ when superimposed on underlying repolarization abnormalities may precipitate lethal arrhythmias.

The setting under which sudden death occurs in narcotic abusers is not amenable to monitoring and therefore to provide

Table VI ECG changes in drug abuse

| | % of patients | Abnormal ECG | QT increased | Brady arrhythmias |
|-----------|---------------|--------------|--------------|-------------------|
| Group I | 34 | 19 | 6 | 11 |
| Group II | 41 | 27 | 14 | 13 |
| Group III | 32 | — | — | 4 |

electrophysiological evidence for the role of ventricular arrhythmias in the acute fatal reactions is not possible. However the existence of an impressive number of repolarization abnormalities and bradyarrhythmias in asymptomatic users of heroin and other abusive drugs makes the possibility of fatal arrhythmias real.

The essential problem revolves around the identification of the subject prone to sudden death. At present longitudinal studies are under way in patients maintained on methadone who are free of other abusive drugs to determine if the initial electrocardiographic abnormalities can revert to normal with time.

Summary

The electrocardiograms (ECG) of 75 asymptomatic individuals admitted to a methadone treatment program were analyzed to determine the effect of street heroin and other drugs of abuse. All patients were free of cardiac renal or pulmonary disease and had no laboratory abnormalities. Two groups of patients were studied. In Group I there were 34 patients on heroin only which was taken within 24 hours of the ECG. Abnormalities were noted in 55 per cent. QT prolongation was found in 19 per cent and significant bradyarrhythmias were found in 19 per cent. In Group II there were 41 patients on methadone with multiple drug abuse. Changes were found in 66 per cent. QT prolongation was noted in 34 per cent, prominent U waves in 32 per cent and bradyarrhythmias were found in 32 per cent.

It is now a well known fact that the sudden death syndrome in addicts may not be a pharmacologic or dosage related phenomenon. The existence of conduction depolarization and repolarization abnor

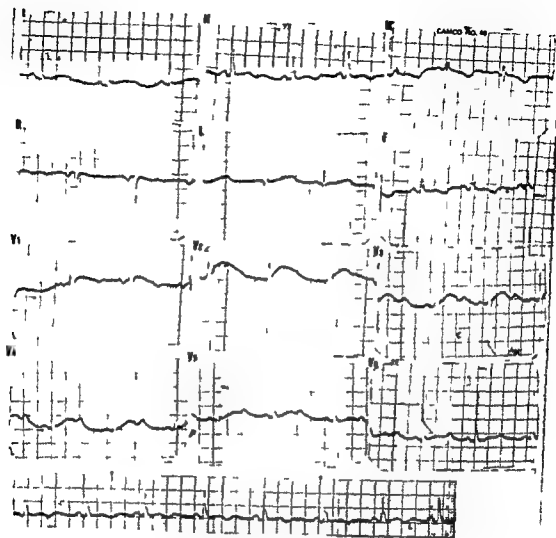


Fig 1 ECG of a 31 year old male drug addict who died suddenly. The QT_c was 0.48 sec and the U wave was prominent.

rate has been found to be variable. Keats and colleagues¹⁰ stressed the tremendous individual variation in response to narcotics and noted variable severity of different side actions in single subjects. Morphine in man has been found by some to slow the heart rate^{11,12} some found an increase^{13,14} and others showed no change.¹⁵ Our findings of bradycardia and tachycardia in a significant proportion of Group I and Group II patients may be related to the variable response to morphine or to yet other related factors or drugs.

Quinine often used as a diluent in illicit heroin preparations alters the ECG in the same manner as quinidine, however a greater dose is required to produce the same effect.¹⁶ Low doses may increase the sinus rate but higher doses produce sinus bradycardia. Its toxic effect may be dose dependent or dose independent. The

rhythm disturbances resulting from quinidine most likely are secondary to a re-entrant mechanism promoted by QT_c prolongation and slow conduction¹⁶ and possibly by an increased dispersion of the refractory period.¹⁷

ECG abnormalities secondary to heroin addiction have been infrequently reported in the literature. Atrial fibrillation has been reported by some.^{18,19} Labl¹⁸ was first to describe proximal atrial fibrillation secondary to heroin intoxication in six patients. In four of these cases, rapid atrial fibrillation with a slow ventricular response was documented prior to therapy with nalorphine hydrochloride. Reversion to regular sinus rhythm after a phase of coarse atrial fibrillation with rapid ventricular response occurred in five. Anoxia and increased vagal tone caused by the drug was postulated as the etiologic mechanism.

Encephalomyocarditis (EMC) virus infection of the mouse aorta*

An ultrastructural study

G E Burch M D

J M Harb Ph D

New Orleans La

The encephalomyocarditis (EMC) virus was first recovered in 1945 from anthropoid apes¹ and later in 1957 from Aotus monkeys.² Since then three fatal outbreaks of myocarditis in domestic swine have been attributed to the EMC virus.³⁻⁵ There are no reports of EMC viral myocarditis in man. However Koch⁶ isolated the virus from earwashings feces blood and cerebrospinal fluid of three patients. A serosurvey of the human populations of southern Louisiana and Peruvian Indians⁷ showed positive neutralizing antibodies to EMC virus in high titer.

In this laboratory EMC virus has been shown to produce myocarditis^{7,8} valvulitis⁹ mural and valvular endocarditis¹⁰ as well as nephritis¹¹ pancreatitis¹² and hepatitis¹³ in experimental animals. This report describes the electron microscopic findings in the aortas of mice infected with EMC virus.

Materials and methods

EMC virus stock was originally obtained from Clasgow.¹⁴ The virus was prepared by

three passages in mouse cardiac cells and 3 passages in L-cells. The titer of the cultivated virus fluid was determined to be 10^6 TCID₅₀. Culture fluid free of virus was used as a control inoculum. Ten one to two-day-old Swiss mice were inoculated intraperitoneally with 0.025 ml of the EMC virus culture fluid and were killed one day later. Two mice were used as controls.

The mice were killed by cervical fracture and the internal organs were quickly removed to expose the aorta. The aorta was bathed *in situ* for 5 minutes with 3 per cent phosphate buffered glutaraldehyde and was then excised and placed in fresh glutaraldehyde for 2 hours. After a 2 to 3 hour phosphate buffer rinse the aorta was post fixed in 1 per cent phosphate buffered osmium tetroxide for 1.5 hours. Dehydration was accomplished in methanol with final embedment in epoxy resin. Thin sections were cut with a Porter Blum MT 2 ultramicrotome or an LKB Ultratome. Sections were stained with 1 per cent aqueous uranyl acetate and lead citrate and

* in the Department of Medicine of the Tulane University School of Medicine and the Charity Hospital of Louisiana at New Orleans, La.

Supported by grant HL-06760 from the National Heart and Lung Institute of the United States Public Health Service. The R. D. Harb was Memorial Fellowship recipient at the Present Heart Laboratory the R. J. Harb and R. E. Harb research in Heart Disease and the F. J. Harb Laboratory.

Received for publication February 19, 1973.

Reprint request to Dr. George E. Burch, 1430 Tulane Avenue, New Orleans, Louisiana 70112.

malities, as well as bradyarrhythmias in a significant number of drug dependent individuals may play a role in the production and facilitation of lethal arrhythmias and may be the mechanism of the acute fatal reaction

REFERENCES

- 1 Helpern M. Fatalities from narcotic addiction in New York City: incidence, circumstances and pathologic findings. *Hum Pathol* 3:13 1972
- 2 Dole V P, Pen W K and Eglitis I. The detection of narcotic drugs, tranquilizers, amphetamines and barbiturates in the urine. *JAMA* 198:348 1966
- 3 Ashman R and Hull E. Essentials of electrocardiography. New York 1937. The Macmillan Company
- 4 Baden M. Narcotic abuse. *N Y State J Med* 72:834 1972
- 5 Steinberg A D and Karliner J S. The clinical spectrum of heroin pulmonary edema. *Arch Intern Med* 122:122 1968
- 6 Silber R and Clarkin E P. Pulmonary edema in acute heroin poisoning. *Am J Med* 27:187 1959
- 7 Helpern M and Rho Y M. Deaths from narcotics in New York City. *N Y State J Med* 66:2391 1966
- 8 Osler W. Edema of left lung morphia poisoning. *Montreal Gen Hosp Rep* 1:291 1880
- 9 Goodman L S and Gilman A. The pharmacologic basis of therapeutics 4th ed. New York 1970. The Macmillan Co. p 238
- 10 Keats A S and Beecher H K. Analgesic potency and side actions. *J Pharmacol Exp Ther* 105:109 1952
- 11 Denton J E and Beecher H K. New analgesics II. A clinical appraisal of the narcotic power of methadone and its isomers. *JAMA* 141:1146 1949
- 12 Wikler A, Fraser H F and Isbell H N. Allyl nomorphine: effect of single doses and precipitation of acute abstinence syndrome during addiction to morphine, methadone or heroin in man. *J Pharmacol Exp Ther* 109:8 1953
- 13 Drew J M, Dripps R D and Comroe J H. Clinical studies on morphine II. The effect of morphine upon the circulation of man and upon the circulatory and respiratory responses to tilting. *Anesthesiology* 7:44 1946
- 14 Eckenhoff J E and Helrich M. Study of narcotics and sedatives for use in preanesthetic medication. *JAMA* 167:115 1958
- 15 Papper E M and Bradley E E. Hemodynamic effects of intravenous morphine and pentothal sodium. *J Pharmacol Exp Ther* 74:319 1942
- 16 Surawicz B and Lasseter K C. Effects of drugs on the electrocardiogram. *Progr Cardiovasc Dis* 13:26 1970
- 17 Han J and Moe G K. Non uniform recovery of excitability in ventricular muscle. *Circ Res* 16:44 1964
- 18 Labl M. Paroxysmal atrial fibrillation in heroin intoxication. *Ann Intern Med* 71:951 1969
- 19 Frand U I, Shim C S and Williams M H Jr. Heroin induced pulmonary edema. *Ann Intern Med* 77:29 1972
- 20 Han J, Millet D, Chizzonitti B and Moe G K. Temporal dispersion of recovery of excitability in atrium and ventricle as a function of heart rate. *Am Heart J* 71:481 1966
- 21 Han J, DeTrazza J, Millet D and Moe G K. Incidence of ectopic beats as a function of basic rate in the ventricle. *Am Heart J* 72:632 1966
- 22 James T N. QT prolongation and sudden death. *Mod Concepts Cardiovasc Dis* 38:35 1969
- 23 Goodman L S and Gilman A. The pharmacologic basis of therapeutics 4th ed. New York 1970. The Macmillan Co. p 106
- 24 Goodman L S and Gilman A. The pharmacologic basis of therapeutics 4th ed. New York 1970. The Macmillan Co. p 206
- 25 Singer K and Lundberg W. Ventricular arrhythmias associated with the ingestion of alcohol. *Ann Intern Med* 77:247 1972



Fig 2 Adventitial cell of aorta from EMC virus infected newborn mouse showing extensive necrosis. The cytoplasm is filled with membrane-bound vesicles (small arrows) and vacuoles (large arrows). Fragmented profiles of rough endoplasmic reticulum (RER) show dilatation and accumulation of dense material within the cisternae. The nuclear chromatin (C) is condensed. Occasional mitochondria (M) are dilated. (Original magnification $\times 31,000$)

age consisted of slight to extensive vesiculation and vacuolization of cytoplasmic membrane systems, dilatation of rough endoplasmic reticulum and accumulation of electron dense material within the cisternae (Fig 2). Often the nuclear chromatin was condensed. Viral crystals were frequently encountered in association with the intracellular necrosis (Figs 3, 4 and 5).

Occasionally the smooth muscle cells in closest proximity to the adventitial tunic were damaged. The necrosis within smooth

muscle cells was identical to that observed in fibroblasts (Figs 4 and 5). Viral crystals also were found in association with the intracellular necrosis in cells which could clearly be identified as smooth muscle cells (Figs 4 and 5).

Viral crystals were found in the aortas of all 10 animals studied. The crystals were composed of dense particles arranged into a cubic configuration (Figs 3 and 4). Viral crystals were located in adventitial fibroblasts and medial smooth muscle cells within fragments of unidentifiable cells



Fig 1 Low magnification electron micrograph of an apparently unaffected portion of aorta (cut in cross section) from ECM viral inoculated newborn mouse. An endothelial cell (*E*) lining the lumen (*L*) is immediately bordered by a band of elastic fibers (*F*₁). Other bands of elastin (*F*₂ to *F*₇) alternate with smooth muscle cells (*SM*) of the media. The adventitial layer (*A*) is adjacent to *F*₇. Cellular components of the adventitia are not clearly demonstrated in this micrograph. A part of a leukocyte (arrow) is visible in the adventitia. (Original magnification $\times 6,900$.)

were examined with a Siemens Elmiskop I electron microscope.

Results

The newborn mouse aorta was roughly separable into three layers (Fig 1). The intima was a single layer of flattened endothelial cells which had an enlarged nucleus that protruded into the aortic lumen. There was no cellular subendothelial layer. The medial tunic was the thickest layer being composed of several layers of smooth muscle cells separated by bands of elastic

fibers (Fig 1). Subjacent to the media was a loosely constructed adventitia, consisting mostly of fibroblastic cells and loose collagen bundles. Occasional blood cells of undetermined types were encountered which possibly had infiltrated into the adventitia.

Cells exhibiting necrosis in the infected animals were found most frequently within the adventitia. These cells were severely damaged and frequently only fragments of cells were found; thus it was difficult to determine the cell types accurately. Dam-



Fig 4 EMC viral infected smooth muscle cell of the media of newborn mouse aorta. A small perinuclear area of intracellular necrosis (top arrow) contains a viral crystal (V). A similar area of necrosis (bottom arrow) is close to the nucleus (N). The smooth muscle cell is bordered on both sides by elastic fibers (F). (Original magnification $\times 19,800$)

and eventually initiate cellular necrosis and aortitis or arteritis which may later develop into scarring of the aorta or arteries. If the vasa vasorum is infected an endarteritis such as is found with syphilis may occur and medial necrosis or inflammation may ensue. Such possibilities need further investigation.

Histologically there has been no vasa vasorum found to penetrate the media of mouse aorta. The structure of the aorta of mouse may be different from that of man. In this study fibroblasts were not found in

the intima and media of mouse aorta. This may have been due to the fact that the mice were killed only a short time after viral inoculation limiting the extent of the disease process. To find a virus injured fibroblastic cell in intima or media after such a short interval of infection may be difficult but it is possible that such cells would be found with advancement of the disease. That the EMC virus can infect fibroblasts of mice is true as discussed above. We suggest that this virus or some other viruses can infect fibroblasts in man.



Fig 3 Adventitial cell of aorta from EMC virus infected newborn mouse. An area of intracellular necrosis (arrow) characterized by membrane bound vesicles and vacuolization is in intimate association with an EMC viral crystal (V). Mitochondria (M) are not swollen. (Original magnification $\times 45,000$)

in the adventitia and free within the extracellular spaces of the adventitia. No viral crystals were found within endothelial cells or within blood cells which were infiltrated into the adventitia.

The aortas of the two control mice were normal.

Discussion

There was direct infection of the adventitial and adjacent smooth muscle layers of the aorta by the EMC virus based on the finding of viral crystals and pathologic changes in the cells of these layers. The crystals had the same appearance as

those previously demonstrated in the myocardium,⁷ heart valves,¹⁰ and kidney¹¹ of infected mice.

Clinically, there are reports of idiopathic medial aortopathy and arteriopathy¹⁵ and nonsyphilitic aortitis¹⁶ in man without a definite etiology or pathogenesis detected. Histologically, fibroblasts are present in all three layers of the aorta of man and the adventitial layer of the vasa vasorum the small nutritional artery of the aorta.¹⁷ Since the EMC virus is highly infectious to fibroblasts in mice it probably can also attack the fibroblasts in the human aorta as well as vasa vasorum or other arteries.

cells of the media in some of the animals. The relationship of these findings to the production of aortitis or arteritis which could also ultimately result in arterio-sclerosis is discussed.

REFERENCES

- 1 Helwig F C and Schmidt E C H Jr Filter passing agent producing interstitial myocarditis in anthropoid apes and small animals Science 102:31 1945
- 2 Roca Garcia M and Sanmartin Barberi C The isolation of encephalomyocarditis virus from Aotus monkeys Am J Trop Med 6 840 1957
- 3 Murnane T G Craighead J E Mondragon H and Shelokov A Fatal disease of swine due to encephalomyocarditis virus Science 132 498 1960
- 4 Gaines J H and Murchison T E Encephalomyocarditis virus infection of swine Vet. Med 56:172 1961
- 5 Koch F Die encephalomyocarditis (EMC) und ihre Abgrenzung von der Poliomyelitis M Kun derheilk. 68:328 1950
- 6 Jonkers A H Serosurvey of encephalomyocarditis virus neutralizing antibodies in southern Louisiana and Peruvian Indian populations Am J Trop Med Hyg 10:593 1961
- 7 Burch G E Harb J M Colcolough H L and Tsui C Y Encephalomyocarditis infection of the newborn mouse myocardium an electron microscopic study Arch. Intern. Med 127:148 1971
- 8 Tsui C Y Burch G E Colcolough H L and Harb J M Early myocardial lesions in encephalomyocarditis (EMC) virus infected mice, Cardiovasc. Res 5:550 1971
- 9 Burch G E and Harb J M Encephalomyocarditis viral valvulitis in new born mouse Experimentia 27:856 1971
- 10 Burch G E Tsui C Y Harb J M and Colcolough H L Mural and valvular endocarditis of mice infected with encephalomyocarditis (EMC) virus Exp Mol Pathol 14:327 1971
- 11 Burch G E Tsui C Y and Harb J M The early renal lesions of mice infected with encephalomyocarditis virus Lab Invest 26:163 1972
- 12 Burch G E Tsui C Y and Harb J M Pancreatic islet cell damage in mice produced by Coxsackie B₁ and encephalomyocarditis viruses Experimentia 28:310 1972
- 13 Harb J M Hiramoto Y and Burch G E Viral hepatitis in encephalomyocarditis virus infected mice (In press)
- 14 Farber P A and Glasgow L A Factors modifying host resistance to virus infection II Enhanced susceptibility of mice to encephalomyocarditis infection during pregnancy Am J Pathol 53:163 1968
- 15 Marquis Y Richardson J B Alexander C K. and Wiggle E D Idiopathic medial aortopathy and arteriopathy Am J Med 44 939 1968
- 16 Restrepo C Trejeda C and Correa D Non syphilitic aortitis Arch Pathol 87:1 1969
- 17 Bloom W and Fawcett D W The textbook of histology 9th ed Philadelphia 1968 W B Saunders Company pp 360-370

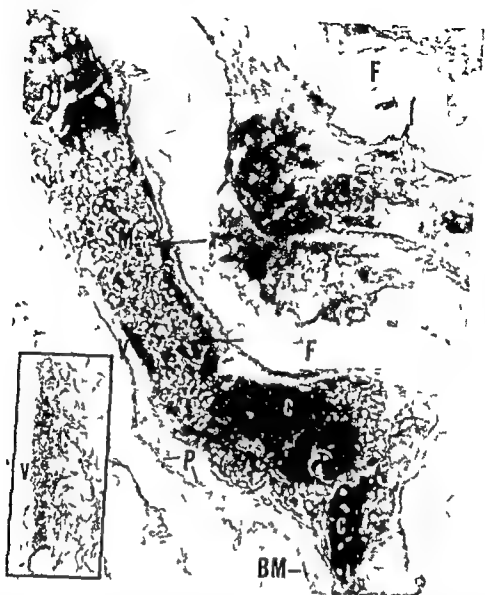


Fig 5 Portions of several smooth muscle cells of the media of a newborn mouse aorta 24 hours following inoculation with EMC viral culture fluid. The central portion of one smooth muscle cell (SM) was filled with densely packed vesicles (arrows). An obliquely sectioned viral crystal (V) (see insert) is present on the outer border of the area of intracellular necrosis. The nuclear chromatin (C) is condensed. Separating the smooth muscle cell layers are elastic fibers (F). Pinocytotic vesicles (P) and an incomplete basement membrane (BM) support the identity of this cell as a smooth muscle cell. (Original magnification $\times 9,800$; insert magnified $\times 34,000$.)

possibly playing a role in the pathogenesis of clinical aortitis or arteritis which could ultimately result in arteriosclerosis. Viral infection of medial smooth muscle cells is evidence that the media can become directly involved in the disease process in addition to being indirectly involved as a result of infiltration of cells which are susceptible to viral infection. Furthermore, intimal involvement could result from migration of virus-injured smooth muscle cells or fibroblasts through fenestrae of the internal elastic lamina. The significance of the aortic damage in relation to scarring

and arteriosclerotic aortic disease remains unknown but deserves investigation.

Summary

Electron microscopic findings in the aortas of 10 mice infected with encephalomyocarditis (EMC) virus are described. Cellular necrosis was found mostly in the adventitia and occasionally in the smooth muscle cells closest to the adventitial tunic. Viral crystals were frequently found in association with the intracellular necrosis. Viral crystals were found in the adventitia of all 10 mice and in the smooth muscle

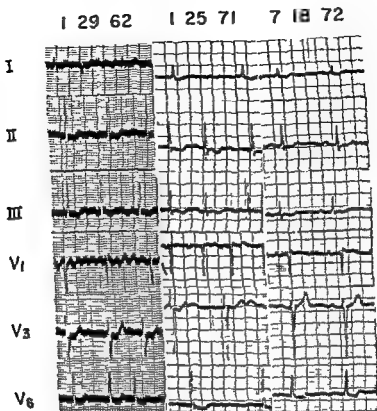


Fig 1 Serial electrocardiogram of Case No. 1 patient H P (No 47 16069)

adequate mitral commissurotomy and it has been maintained for one year.

Case 2 L G a Caucasian woman developed rheumatic fever at age 5 years. A heart murmur was noted at age 29 and digitoxin 0.1 mg daily was started. Later he developed bronchial asthma and the cardiac functional state gradually declined. Atrial fibrillation developed at age 47 when she was first seen at the University Hospitals. In addition to bronchial asthma she had a mitral facies, a precordial lift, cardiomegaly and the murmurs of mitral stenosis and mild mitral and aortic insufficiency. The electrocardiogram showed atrial fibrillation with fibrillatory waves 3 mm in amplitude in Lead V. A closed mitral commissurotomy was done in the same year with temporary improvement in the cardiac functional state. Diuretic therapy was added to digitalis at age 55 but wheezing and dyspnea became almost constant requiring her to sleep on four pillows. At age 56 prednisone therapy was begun which afforded her considerable relief from the asthma. Coarse atrial fibrillation was noted on every re-examination for 16 consecutive visits until age 59 when it was found that sinus rhythm had returned spontaneously. The electrocardiogram at this time showed right ventricular hypertrophy, left atrial disease, short runs of atrial tachycardia with long P axis and 1st degree A V block (Fig 2).

At age 60 diabetes mellitus was discovered. She was last seen in 1972 at age 61 and had isorhythmic A V dissociation with a sinus mechanism and her major problem then was osteoporosis. Atrial fibrillation which had persisted for a total of twelve years after an adequate mitral commissurotomy converted spontaneously to sinus rhythm which has been maintained for the last two years.

Case 3 A C a Caucasian woman had acute arthritis at age 8 and 15 years and at age 24 a cardiac murmur was noted. At age 55 an electrocardiogram showed sinus rhythm with frequent premature atrial contractions. During the following five years she experienced frequent paroxysms of a tachyarrhythmia, recurrent lung infections with dyspnea and mild blood streaking of the sputum and gradual reduction in exercise tolerance. Nocturnal dyspnea, pulmonary congestion and atrial fibrillation appeared at age 60 during a severe respiratory infection. Examination revealed a mitral facies, right ventricular lift, loud S₁ and P₂ and a long diastolic apical rumble. Quinidine therapy restored sinus rhythm. She had a closed mitral commissurotomy in the same year (age 60) and atrial fibrillation recurred twice within two weeks and once in the following year. Digitalis and quinidine restored sinus rhythm each time. Quinidine had to be stopped at age 61 because of cinchonism and coarse

Spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and rheumatic mitral valve disease

Description of three patients

Thomas J Zimmerman, M D
Lotfy L Basta, M D, M R C P
Lewis E January, M D
Iowa City Iowa

Atrial fibrillation is a complication of rheumatic mitral disease probably determined less by the severity of the valve lesion than by the age of the patient.^{1,2} The majority of patients with mitral valve disease eventually develop atrial fibrillation coincident with clinical deterioration³ and often it may be paroxysmal initially before it becomes the established rhythm.⁴ Once established for a few years, attempts at cardioversion may either fail to restore sinus rhythm^{5,6} or if successful, sinus rhythm rarely is maintained for more than a few weeks or months.^{7,12}

That patients with mitral valve disease who have had atrial fibrillation for many years, occasionally may revert spontaneously to sinus rhythm and maintain that rhythm has not been previously emphasized. One of the authors (L E J) has personally followed 120 patients for up to twenty years after closed commissurotomy. Three of these patients spontaneously reverted to sinus rhythm after having atrial fibrillation for over ten years, they are the subject for this report.

Case reports

Case 1 H P, a Caucasian woman developed pneumonia at age 18 years when a heart murmur was first detected. Pneumonia recurred with her first pregnancy three years later. At age 31 she was seen at the University Hospitals with dyspnea, coughing and wheezing and the findings of a right ventricular systolic lift, loud S₁ and an apical diastolic rumble. Chest x ray showed pulmonary venous congestion and a prominent left atrium and right ventricle. Pneumonia recurred at age 34 and the following year she developed congestive heart failure during treatment for leg burns. Digitalis and diuretics were given and later discontinued after five months. Atrial fibrillation developed at age 36 accompanied by an increase in pulmonary congestion. Digitalis improved her symptoms but dyspnea on mild exertion and atrial fibrillation continued. A closed mitral commissurotomy was performed at age 37 and a left atrial appendage thrombus was removed. She developed an expressive aphasia which subsequently cleared. Dyspnea improved and only minimal mitral insufficiency was noted. Atrial fibrillation was found on eight subsequent yearly visits. Initially the fibrillation waves were 2 to 3 mm in amplitude in Lead V₁ but at age 47 they became 1 mm in amplitude in the same lead (Fig. 1). Sinus rhythm with first degree A V block spontaneously returned at age 47 and has been maintained until the present time. Sinus rhythm returned 12 years following the onset of atrial fibrillation and 10 years after an

From the Cardiovascular Division, University of Iowa and Veterans Hospitals, Iowa City, Iowa.
Supported by Program Project Grant No. HL14388 and by Iowa Heart Association Grant No. 05467.

Received for publication Jan. 2, 1973.

Reprint requests to Lotfy L. Basta, M.D., M.R.C.P., Department of Internal Medicine, University Hospital, Iowa City, Iowa 52242.

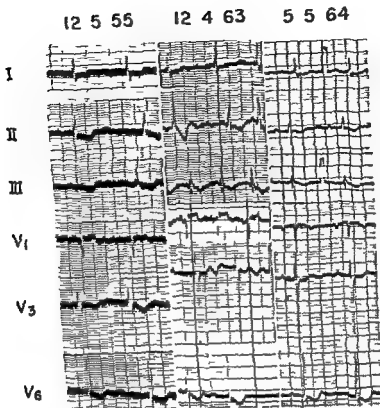


Fig 3 Serial electrocardiogram of Case No 3 patient A C (No 53-13783)

cardioverted or sinus rhythm was recovered only transiently. Twenty two of 25 patients described by Semer and associates⁷ with more than five years atrial fibrillation did not maintain sinus rhythm for six months following electroversion. Similarly no patient with atrial fibrillation of more than five years maintained sinus rhythm for one year following cardioversion in the series reported by Radford and associates,⁸ Bailey and colleagues,⁹ Olesen and co workers,¹⁰ or Upton and Honey.¹¹ In our three patients with atrial fibrillation for more than ten years one would have expected that attempts at defibrillation either would have failed or if successful that sinus rhythm could not be maintained. Instead sinus rhythm spontaneously returned.

In patients with rheumatic heart disease established atrial fibrillation has been correlated with pathologic changes in the atrial wall^{12,13} and particularly with the extent of disease of the sinoatrial node and internodal pathways.^{14,15} Moreover there is recent evidence suggesting that the persistence of atrial fibrillation accentuates

pathologic damage of the left atrium probably through dilatation and increased wall tension.¹⁶ This is generally thought to further decrease the chances of recovery of sinus rhythm. On the other hand Holzmänn¹⁷ described one case of atrial fibrillation with spontaneous reconversion to sinus rhythm in whom the left atrium was found at autopsy to be converted into a fibrous sac. The author suggested that initially the atrial fibrillation was triggered by left atrial involvement and reconversion occurred when all of the left atrial myocardium was destroyed.

Recent atrial fibrillation usually shows coarse f waves in the electrocardiogram¹⁸ but fibrillation waves become finer with chronicity¹⁹ and with digitalis treatment.²⁰ Low amplitude f waves in the electrocardiogram generally are regarded as an unfavorable sign when defibrillation is to be considered.²¹ It is remarkable that the three patients included in this report maintained the coarse fibrillatory waves throughout the years and in spite of digitalis therapy a finding that may have been

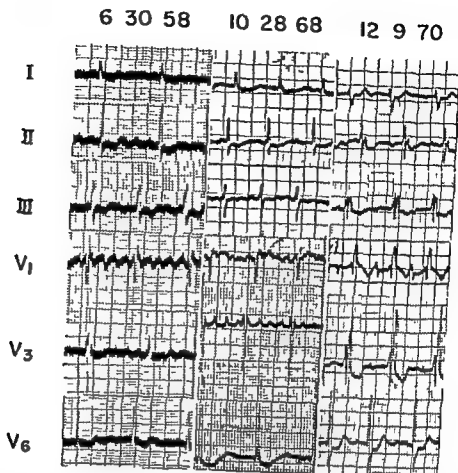


Fig 2 Serial electrocardiogram of Case No 2 patient L. G. (No 58 11748)

atrial fibrillation became the established heart rhythm noted on all subsequent visits for the next eight years. She developed an anterior myocardial infarction at age 67. At age 70 intermittent atrial flutter was noted on one electrocardiogram (Fig 3). Sinus rhythm spontaneously appeared at age 71 with 1st degree A-V block. Digoxin had been her only medication for years. She developed heart failure at age 74 and a cerebral vascular stroke at age 75 causing her death while still in sinus rhythm. Thus this patient had atrial fibrillation which was recurrent for two years and established for nine years. It spontaneously reverted to sinus rhythm eleven years after a closed mitral commissurotomy and the normal rhythm persisted for four years thereafter.

Discussion

We have described three patients representative of a rarely recognized phenomenon^{11,12} that is difficult to explain from our present knowledge of the pathogenesis and course of atrial fibrillation in rheumatic heart disease. It is possible, however, that this phenomenon is not this rare, and many more cases will be recognized when adequately looked for.

Soon after successful mitral valve sur-

gery, patients with atrial fibrillation may return to sinus rhythm occasionally spontaneously but more commonly with therapeutic defibrillation.^{4,12} Successful surgery may enhance the resumption of sinus rhythm, probably as a result of atrial decompression.¹¹ It is conceivable that in our three patients the favorable effects of surgery may have allowed significant sustained reduction in left atrial pressure and wall tension with subsequent recovery of the sinus rhythm after more than ten years. This assumption is however challenged by the observation that such reversion may still occur when there is almost total damage of the left atrium as determined pathologically.¹¹

It has been shown that the duration of atrial fibrillation is the most important single determinant of the immediate success and long term maintenance of sinus rhythm following cardioversion in patients with rheumatic mitral disease. In the series reported by Lown,⁶ 12 of 18 patients with atrial fibrillation of more than ten years duration either could not be electrically

Mitral disc variance (Harken prosthesis)

Sukh Dev Sharma M D
Robert M Easley Jr M D
Lawrence I Zaroff M D
Sidney Goldstein M D
Rochester N Y

Although the operative mortality and morbidity rate in surgical replacement of cardiac valves has decreased considerably in the past decade the development of mechanical dysfunction of prosthetic valves continues to be a serious problem. Variance with ball valve prostheses in both aortic and mitral positions is well documented.¹⁻⁶ Low profile discoid valve prostheses have been used with increasing frequency for mitral valve replacement because of certain apparent advantages they have over ball valve prostheses in this position being light in weight having less inertia and having a short cage which occupies less space in the ventricle.⁷⁻¹¹ Nevertheless malfunction of various types of these disc prostheses has also been reported.¹²⁻²² We report the first such variance in a Harken Surgitool Discoid mitral valve prosthesis and discuss some of the unique features of disc variance of the mitral valve.

Case report

Patient A N a 31 year old housewife with a history of rheumatic fever in childhood had been asymptomatic until April 1968 when during her

fourth month of pregnancy she developed severe exertional and paroxysmal nocturnal dyspnea and orthopnea. She was hospitalized in acute pulmonary edema and was found to have rheumatic heart disease with mitral stenosis and aortic regurgitation (N Y H A Class IV). She was treated with digitalis and diuretics and improved. Subsequently she delivered a full term infant without further difficulty. Over the next 1½ years she developed progressive exertional dyspnea. In June 1970 she was readmitted for cardiac catheterization because of limiting symptoms (N Y H A Class III). Severe mitral stenosis (calculated mitral valve area 0.9 cm²) and minimal aortic regurgitation were demonstrated (Table I). Mitral valve replacement with a small (S/M 432 H) Harken Mitral Discoid valve prosthesis Magovern series (turtle neck) was performed on June 11 1970. Her postoperative course was uneventful and she returned to work as a nurse's aide. She continued to take digoxin warfarin and prophylactic penicillin and was asymptomatic until 23 months after surgery when symptoms of palpitation progressive dyspnea orthopnea and paroxysmal nocturnal dyspnea suddenly reappeared. After 10 days she sought medical attention and was hospitalized on May 4 1972. There was no history of fever hemoptysis or chest pain. On examination her pulse was 100 per minute and regular. Blood pressure was 90/48 mm Hg. The jugular venous pressure was not elevated. Basilar rales were present bilaterally. The apical impulse was displaced slightly to the left and a right ventricular lift was palpable. An intermittent Grade III/VI pansystolic murmur radiating toward the axilla and a Grade II/VI mid-diastolic

From the Department of Medicine and Surgery, Rochester General Hospital and University of Rochester, School of Medicine and Dentistry, Rochester, N. Y.
D. Sharma was supported by a Research Fellowship in the Genesee Valley Heart Association.
Received for publication April 14 1973.
Reprint requests to: Robert M. Easley, Jr., M.D., Rochester General Hospital, 1425 Highland Ave., Rochester, N. Y. 14621.

relevant to their subsequent spontaneous reversion to sinus rhythm

It is interesting to note that with return of sinus rhythm the electrocardiograms consistently showed impaired atrioventricular conduction, a phenomenon that has been previously noted with therapeutic cardioversion of atrial fibrillation.²⁴ It is also notable that with return of sinus rhythm there was practically no change in the clinical state or the digitalis requirements in any of these patients

With modern treatment and particularly with improvement of surgical techniques many more patients with rheumatic mitral disease may survive until old age. Perhaps return of sinus rhythm after many years of established atrial fibrillation may not be too uncommon in such patients. Further observation should determine the frequency of this phenomenon

Summary

Three patients with rheumatic mitral stenosis had had atrial fibrillation for over ten years before spontaneously restoring normal sinus rhythm. Each had undergone a closed mitral commissurotomy ten to twelve years before the conversion. Their ages at the time of resuming normal rhythm were 47, 59, and 71 years. This unusual phenomenon has no satisfactory explanation by our present knowledge of the natural history of mitral valve disease and the pathogenesis of atrial fibrillation.

The authors wish to acknowledge the valuable secretarial help of Sara Fraleigh

REFERENCES

- 1 Wood P. An appreciation of mitral stenosis. *Br Med J* 1:1051 1954
- 2 Rowe J, C Bland E, F Sprague H II and White P D. The course of mitral stenosis without surgery: ten and twenty year perspectives. *Ann Intern Med* 52:741 1960
- 3 Olesen K H. The natural history of 271 patients with mitral stenosis under medical treatment. *Br Heart J* 24:349 1962
- 4 Selzer A and Cohn K E. Natural history of mitral stenosis—a review. *Circulation* 45:878 1972
- 5 Bell H, Pugh D and Dunn M. Failure of cardioversion in mitral valve disease. *Arch Intern Med* 119:257 1967
- 6 Lown B. Electrical reversion of cardiac arrhythmia. *Br Heart J* 29:469 1967
- 7 Semer H, Hultgren H, Kleiger R and Braniff B. Cardioversion following prosthetic mitral valve replacement. *Circulation* 35:523 1967
- 8 Radford M D and Evans D W. Long term results of D C reversion of atrial fibrillation. *Br Heart J* 30:91 1968
- 9 Bailey G W H, Braniff B A, Hancock E W and Cohn K E. Relation of left atrial pathology to atrial fibrillation in mitral valvular disease. *Ann Intern Med* 69:13 1968
- 10 Olesen K H, Andersen M, Flensstedt Jensen E, Hansen J F, Steinness E and Winkel P. D C conversion of atrial fibrillation in mitral stenosis. In: Symposium on Cardiac Arrhythmias. Elsinore, Denmark. Södertälje, Sweden 1970. A B Astra p 393
- 11 Upton A R M and Honey M. Electroconversion of atrial fibrillation after mitral valvotomy. *Br Heart J* 33:732 1971
- 12 Warris E, Kreis K E and Salokannel M. Factors influencing persistence of sinus rhythm after D C shock treatment of atrial fibrillation. *Acta Med Scand* 189:161 1971
- 13 Holzmänn M. Basic mechanism of atrial fibrillation. In: Symposium on Cardiac Arrhythmias. Elsinore, Denmark 1970. Södertälje, Sweden 1970. A B Astra p 92
- 14 White P D. Basic mechanism of atrial fibrillation. In: Symposium on Cardiac Arrhythmias. Elsinore, Denmark. Södertälje, Sweden 1970. A B Astra p 92
- 15 Fisher R D, Mason D T and Morrow A G. Restoration of sinus rhythm after mitral valve replacement. *Circulation* 37:173 1968
- 16 Rossi L. Histopathologic features of cardiac arrhythmias. Milano 1969. Casa Editrice Ambrosiana page 110
- 17 Hildebrand H E. Der Sinusknoten bei arrhythmie. *Virchows Arch (Pathol Anat)* 337:1515 1964
- 18 Hudson R E B. The human pacemaker and its pathology. *Br Heart J* 22:153 1960
- 19 Sims B A. Pathogenesis of atrial arrhythmias. *Br Heart J* 34:336 1972
- 20 Davies M J and Pomerance A. Pathology of atrial fibrillation in man. *Br Heart J* 34:520 1972
- 21 Aberg H. Coarse and fine atrial fibrillation: method of evaluation, relation to mechanism and aetiology of fibrillation. Symposium on Cardiac Arrhythmias. Elsinore, Denmark. Södertälje, Sweden 1970. A B Astra p 43
- 22 White P D. F wave size, influence of time and treatment. Symposium on Cardiac Arrhythmias. Elsinore, Denmark. Södertälje, Sweden 1970. A B Astra p 51
- 23 Stern S F. Wave size correlation to result of cardioversion. Symposium on Cardiac Arrhythmias. Elsinore, Denmark. Södertälje, Sweden 1970. A B Astra p 51
- 24 Lown B, Perloff M G, Kaidbey S, Abe T and Harken D E. Cardioversion of atrial fibrillation: A report on the treatment of episodes in 50 patients. *N Engl J Med* 269:325 1963

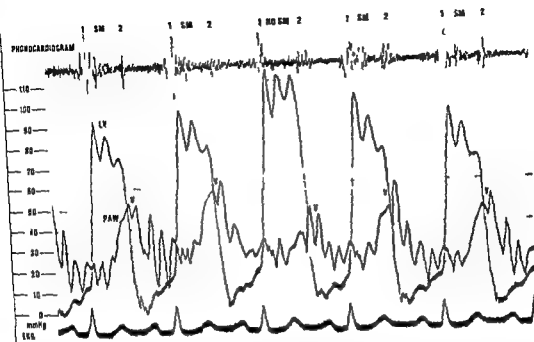


Fig 2 Simultaneous phonocardiographic (at the apex) left ventricular pressure pulmonary artery wedge pressure and electrocardiographic tracings Note that left ventricular systolic pressure and PA wedge V wave are variable from beat to beat The third cycle shows better valve function with lower V wave higher LV systolic pressure and absence of systolic murmur of mitral regurgitation LV = left ventricular pressure PAW = pulmonary artery wedge pressure SM = systolic murmur V = V wave of PAW

Table 1 Serial hemodynamic data

| | 6 3 70 | | 5 3 72 |
|------------------------------|--------------------------|--------------------------|--------------------------|
| | Resting | Exercise | Resting |
| Right atrium | 3 mm Hg | | 7 mm Hg |
| Right ventricle | 46/7 mm Hg | | 55/8 mm Hg |
| Pulmonary artery | 46/22(30) mm Hg | 100/58(75) mm Hg | 55/26(42) mm Hg |
| Pulmonary artery wedge | (23) mm Hg | (60) mm Hg | (35) mm Hg |
| Left ventricle | 100/4 mm Hg | 135/0 mm Hg | 100/16 mm Hg |
| Mitral valve gradient | 26 mm Hg | 59 mm | 18 mm Hg |
| Calculated mitral valve area | 0.9 cm ² | 1.1 cm ² | 1.1 cm ² |
| Cardiac index | 3.0 L/min/M ² | 5.1 L/min/M ² | 2.4 L/min/M ² |

the mitral valve because of several apparent theoretical disadvantages inherent with the ball valve prosthesis in this position^{1,2,9} These include protrusion of bulky valve structure into the left ventricular outflow tract contributing to low cardiac output arrhythmias due to impingement on the interventricular septum partial obstruction to blood flow by restricted orifice size and relatively slow response of the ball due to

static inertia Matloff and colleagues¹¹ reported striking hemodynamic improvement and fall in pulmonary vascular resistance in the early postoperative periods in patients with Harken Discoid mitral prostheses Nevertheless both the complications of thrombosis and variance have been reported with mitral disc prostheses as well and can be responsible for serious congestive heart failure^{9,11,22}



Fig 1. A through C Frames from 70 mm fluoroscopic spot films A abnormal wear of the superior edge of the disc is seen B and C worn disc is seen incompletely covering the inflow orifice (B) and projecting into the left atrium (C)

murmur at the apex and an early decrescendo Grade II/VI diastolic murmur in the left parasternal region were heard. The opening and closing disc sounds varied in intensity from beat to beat. At times the opening click was inaudible and the second sound opening click interval was variable. There was no hepatomegaly or peripheral edema. The electrocardiogram revealed normal sinus rhythm, probable left atrial enlargement, and non specific ST segment

and T wave abnormalities. The chest roentgenogram as compared to the most recent x rays showed left atrial and left ventricular enlargement, interstitial edema, distended upper lobe veins, and interlobar fluid in the minor fissure. For the evaluation of function of the mitral valve prosthesis, 70 mm fluoroscopic spot films were taken. Distortion, loss of spherical shape, and discontinuity of the superior edge were noticed. In systole, the disc usually was seated at an abnormal angle with the superior aspect projecting through the ring of the cage toward the left atrial cavity (Fig 1). The diagnosis of variance of the mitral disc prosthesis was made. Cardiac catheterization performed on May 5, 1972, demonstrated a mitral valve gradient, left atrial V wave, and left ventricular pressures which varied from beat to beat depending upon the degree of prosthetic malfunction (Fig 2). Mild aortic regurgitation was reconfirmed. The left ventricular angiogram showed abnormal motion of prosthetic valve disc and severe mitral regurgitation. Immediately following the catheterization, reoperation was carried out. At surgery, the prosthesis was examined and the disc was found to be markedly deficient in substance (Fig 3). It easily lifted out of the cage and was approximately half of its expected size. A Starr-Edwards ball valve prosthesis (Titanium ball and cloth covered ring Model 6320T) was inserted. During surgery, the patient developed complete AV block and required ventricular pacing. This problem persisted postoperatively and a permanent transvenous Medtronic Demand pacemaker was implanted on May 15, 1972. Following an otherwise uneventful postoperative course, the patient was discharged on digoxin and warfarin. She is asymptomatic at the time of this writing.

The pathologic examination of the prosthesis showed no adherent fibrosis or clot around the disc. The diameter of the disc measured 2.6 cm, as compared to 2.79 cm of the normal.

Discussion

The use of intracardiac valvular prostheses during the last decade has prolonged survival and improved the clinical state of patients with advanced valvular heart disease. However, various late complications such as embolism, hemolysis, periprosthetic infection, detachment of the sewing ring, and thrombosis of the prosthesis are problems faced by these patients after surviving valve replacement. Intrinsic ball valve malfunction¹⁻⁴ due to degeneration of the ball²³ causing swelling,²⁴ impaction,²⁴⁻²⁵ fissuring, fracture or fragmentation²³ leading to dislodgement²⁶ and embolization²³ have been reported with aortic and mitral prostheses of Starr-Edwards, Smeloff-Cutter, SCDK Cutter, and Magovern-Cromie.

Low profile discoid prostheses have been used with increasing frequency to replace

been reported more frequently in the mitral position than in the aortic.^{1, 11, 15} With the development of new valves with silastic occluders ball variance has been only occasionally reported with series 1200 Starr Edwards aortic and other prosthetic valves.⁶ Abnormalities in the disc mitral valve have been observed in Hufnagel,^{1, 12} Kay Shiley,^{14, 15} Kay Suzuki Cross Jones,¹⁷ Beall Surgitool^{20, 21} and Hammer Smith²² valves. In our experience with 60 patients with Harken Surgitool Discoid prostheses this is the only patient who has developed variance. Disc variance secondary to surface abrasion occurs because of wear of the disc edges producing insufficiency of the prosthesis. This may vary from beat to beat depending on the disc position across the inflow orifice as was seen in our patient. Massive mitral incompetence has also been observed secondary to cocking of the disc.¹⁶ Edgett and associates¹⁸ reported intermittent mitral insufficiency in four patients produced by an aortic regurgitant jet causing tilting of the prosthetic disc.

Whatever the cause of mechanical failure the morbidity and mortality rate associated with prosthetic valve dysfunction necessitates early surgical intervention even though reoperation is technically more difficult and carries a higher surgical risk. An operative mortality rate of 13 per cent for patients with ball variance has been reported by Hylen and colleagues.¹ Since the first successful replacement of a thrombosed mitral ball valve prosthesis in 1965 by Spencer and co-workers,²³ reoperation for replacement of the valve prosthesis has been the treatment of choice in patients with mitral prosthetic dysfunction. Herr and associates⁴ go a step further in advocating replacement even in asymptomatic patients having evidence of prosthetic malfunction by auscultation and phonocardiography. Therefore it seems justified to recommend frequent re-examination including phonocardiographic, fluoroscopic and where possible echocardiographic evaluation of all patients with mitral prostheses. Whenever there is an acute alteration in clinical status of the patient in the post-operative phase complete restudy is imperative. The presence of asynchrony, absence or diminution of prosthetic sounds or

appearance of a new murmur persistent or transient should alert a physician to the possibility of ball or disc variance. Early recognition of this complication is essential for successful surgical treatment.

Summary

A case of variance of a Harken Surgitool Discoid mitral prosthesis is reported. Prosthetic malfunction was suspected because of sudden deterioration of cardiac status after a previously stable postoperative course. The appearance of new murmurs as well as variable intensity and timing of the prosthetic sounds were noted in this patient. The diagnosis was confirmed radiologically and by cardiac catheterization. The patient underwent successful replacement of the mitral prosthesis.

The problem of prosthetic variance is discussed with emphasis on diagnostic findings in malfunction of mitral prosthesis.

REFERENCES

1. Hylen J C. Mechanical malfunction and thrombosis of prosthetic heart valves. *Am J Cardiol* 30:396 1972.
2. Hylen J C. Durability of prosthetic heart valves. *AM HEART J* 81:299 1971.
3. Hylen J C, Kloster F E, Starr A, and Griswold H E. Aortic ball variance: diagnosis and treatment. *Ann Intern Med* 72:1 1970.
4. Herr H H, Kloster F E, Yukiyasu S, and Starr A. Diagnosis and management of ball variance following aortic valve replacement. *Circulation* 35 (Suppl 11):141 1968.
5. Starr A, Rodney H H, and Wood A. Mitral replacement: review of six years experience. *J Thorac Cardiovasc Surg* 54:331 1967.
6. Bigelow J C, Herr R H, Wood J G, and Starr A. Multiple valve replacement: review of five years experience. *Circulation* 38:656 1968.
7. Beall A C, Bloodwell E D, Bricker D L, Okes J E, Cooley D A, and DeBakey M E. Prosthetic replacement of cardiac valves. *Am J Cardiol* 23:250 1969.
8. Behrendt D M, and Austen W G. Current status of prosthetics for heart valve replacement. *Progr Cardiovasc Dis* 15:No 4 1973.
9. Vasko J S, and Leighton R F. Acute massive mitral regurgitation resulting from disc valve malfunction. *Ann Thorac Surg* 6:564 1968.
10. Dalen J E, Matloff J M, Evans C L, et al. Early reduction of pulmonary vascular resistance after mitral valve replacement. *N Engl J Med* 277:387 1967.
11. Matloff J M, Dalen J E, Dexter L, et al. Hemodynamic response to discoid mitral valve

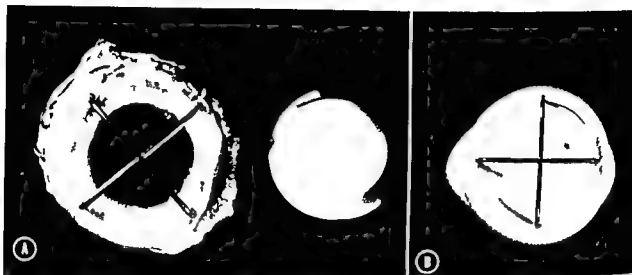


Fig 3 A and B Appearance of prosthetic valves A abnormally worn Harken Discoid mitral prosthesis after removal from this patient B normal prosthesis is shown for comparison

Sudden deterioration in the cardiovascular status in a patient who has been doing well in the postoperative period is a common mode of clinical presentation. Acute massive mitral regurgitation resulting from disc valve malfunction has been reported.¹⁻¹⁶ The early recognition of ball as well as disc variance by auscultation and phonocardiography has been well described and emphasized.^{1-4, 10, 20, 22}

Total disappearance or a decrease in the intensity of aortic opening sound as compared to the closing sound is diagnostic of aortic prosthetic ball variance.^{1-4, 20} The presence of variable intensity of the mitral prosthetic opening and closing clicks, diminution or complete disappearance of the opening click, and delayed and variable aortic closure-opening click interval, are regarded as characteristic features of mitral prosthetic variance.¹¹⁻¹³ The appearance of new cardiac murmurs, constant or intermittent, suggests prosthetic malfunction. Hylen and colleagues²⁴ described sound spectrography in the detection of prosthetic dysfunction associated with aortic valve prosthesis and found abnormal sound spectrogram in 78 per cent of patients with ball variance. Pfeifer and associates²⁰ described periodic delayed opening of the mitral valve prosthesis associated with abnormal motion of the cage by echocardiography.

Radiological examination of the mitral prosthesis in patients thought to have prosthetic valve dysfunction is a very important diagnostic tool. Lansing²⁵ reported a double

exposure appearance of the prosthesis in a prolonged exposure chest film as a common sign of loose mitral valve prosthesis. In our patient not only malposition but change in the shape of the disc could be determined on 70 mm cut film x rays.

The value of cardiac catheterization to confirm and establish the degree of prosthetic malfunction cannot be overemphasized. However, it is necessary to be aware of factitious mitral insufficiency or even stenosis²⁶ caused by inadvertent passage of the cardiac catheter through the struts of the prosthesis when passed retrograde into the left ventricle. This may temporarily interfere with motion of the ball or disc.

Malfunction of mitral ball as well as disc prostheses due to thrombotic complications causing either impairment of ball movement in the cage^{11, 20, 27} or repeated embolic episodes²⁴ has been reported. Swelling of the ball resulting in critical obstruction to flow through the prosthesis has also been described.²⁴ This has been considered to be due to degeneration of the poppet secondary to the presence of some chemical impurities in its basic material. Absorption of blood lipids²⁸ as well as certain drugs and chemical compounds²⁴ by a silicon ball leading to its degeneration has been shown by drug diffusion studies and chemical analysis of poppets. These problems have been more frequent with series 1000 Starr Edwards aortic prosthesis than in the Starr Edwards mitral valve.^{1, 2, 6, 27} In contrast with Smeloff Cutter prosthesis ball variance has

Nonpenetrating cardiac injuries A collective review

A James Liedtke MD
William E DeMuth Jr MD
Hershey Pa

Violent injury now accounts for the majority of deaths among persons below the age of 40 years and as described recently by the National Research Council of the National Academy of Sciences represents the neglected epidemic of modern society.¹ In a consideration of the sequelae of injury to various organ systems cardiac damage is particularly hazardous and accounts for one of the leading causes of death in traumatic injury. However its diagnosis is often difficult and as recently stated by the medical examiner for the City of Philadelphia represents the commonest unsuspected visceral injury responsible for death in fatally injured accident victims in that city.² If this estimate is correct then presumably the incidence of heart damage may be much higher in rural areas where deceleration injuries account for much larger proportions of death.³

Whereas penetrating wounds to the chest involving the heart are seldom unsuspected by virtue of the nature of the wounds nonpenetrating chest trauma injuring the heart is often found in combinations with other organ system injuries including major fractures ruptured viscera and other less serious but more noticeable injuries all of which tend to overshadow and obscure the cardiac damage. A recent review by Parmley and colleagues⁴ demonstrates the infrequency with which the diagnosis is made

In 353 instances of myocardial rupture secondary to nonpenetrating trauma discovered at autopsy a cardiac lesion was suspected only once prior to death. Quite obviously injuries associated with less hemodynamic derangement are even more likely to be overlooked by clinicians.

Although young adults are most frequently the victims of nonpenetrating heart injury all age groups are susceptible. Cardiac injury has been described in the in utero fetus due to maternal injury⁵ during labor and delivery^{6,7} and in very early childhood.⁸

Although nonpenetrating heart injuries have been described in sporadic case reports for hundreds of years Beck⁹ was the first to systematically document this entity by his clinical and laboratory observations in 1935. It is the purpose of the present communication to extend these observations first by reviewing the several types of cardiac lesions found in nonpenetrating trauma second by presenting more recent data relating pertinent clinical manifestations and diagnostic clues and lastly by discussing currently available information concerning therapy.

Mechanisms of injury

Before the days of high speed automobile travel nonpenetrating chest trauma with subsequent heart damage was quite un-

From the Department of Medicine and Surgery, College of Medicine, Pennsylvania State University, The Milton S. Eshley Medical Center, Hershey, Pa.
Received for publication Jan. 15, 1973.
Reprint requests: William E. DeMuth, Jr., M.D., Department of Surgery, The Milton S. Eshley Medical Center, Hershey, Pa. 17033.

- replacement *Circulation* 37 (Suppl II) 96 1968
- 12 Vogel J H K Paton B C Ovary H R et al Abnormal hemodynamic function after disc mitral valve replacement *Circulation* 39 (Suppl I) 141, 1969
- 13 Hopeman A R Treasure R L and Hall R J Mechanical dysfunction in caged lensa prosthesis *J Thorac Cardiovasc Surg* 60:51 1973
- 14 Paton B C Reoperation of fibrinous stenosis of disc mitral valve prosthesis *J Thorac Cardiovasc Surg* 57:726 1969
- 15 Paton B C, Vogel J H K, Ovary H et al Follow up results on Kay Shiley valve in Brewer L A ed *Prosthetic heart valves* Springfield Ill 1969 Charles C Thomas Publisher
- 16 Low H B C and Lefemine A A Acute mitral insufficiency due to jamming of a disc valve prosthesis *Annals of Thoracic Surgery* 4:71 73 1967
- 17 Braunwald N S and Detmer D E Critical analysis of status of prosthetic valves and homografts *Progr Cardiovasc Dis* 11:113 1968
- 18 Edgett J W Nelson W P Hall R J et al A complication of valve replacement by a caged lensa prosthesis *Circulation* 36:422 1967
- 19 Cross F S Akao M and Jones, R D The evaluation of experimental mitral valve prosthesis in the dog *Surgery* 65 89 1969
- 20 Javier R P Hildner F J Berry W et al Systemic embolization and the Beall mitral valve prosthesis *Ann Thorac Surg* 10:20 1970
- 21 Beall A C Bloodwell R D Arbeagast N R et al Mitral valve replacement with dacron covered disc prosthesis to prevent thromboembolism Clinical experience in 202 cases in Brewer L A ed *Prosthetic heart valves* Springfield Ill 1969 Charles C Thomas Publisher pp 319 325
- 22 Samaan H A Acute massive mitral regurgitation resulting from disc valve replacement of mitral valve *J Cardiovasc Surg* 10:477 1969
- 23 Roberts W C and Morrow A G Fatal degeneration of the silicone rubber ball of the Starr Edwards prosthetic aortic valve *Am J Cardiol* 22 619 1968
- 24 Leatherman L L Leachman R D McConn R G, et al Malfunction of the mitral ball valve prosthesis due to swollen poppet *J Thorac Cardiovasc Surg* 57 160 1969
- 25 Lee S J K Zaragoza A J Callaghan J C et al Malfunction of mitral valve prosthesis (Cutter Smeloff) *Circulation* 41 479 1970
- 26 Krosnick F E Death due to migration of the ball from an aortic valve prosthesis *JAMA* 191 105 1965
- 27 Starr A Mitral valve replacement with ball valve prosthesis *Br Heart J* 33 (Suppl) 47 1971
- 28 Hylen J C Kloster I E Herr R H et al Phonocardiographic diagnosis of aortic ball variance *Circulation* 38:90 1969
- 29 Sanderson R G Hall A D and Thomas A N The clinical diagnosis of ball variance in a mitral valve prosthesis *Ann Thorac Surg* 6:473 1968
- 30 Pfeifer J Goldschlager N Sweatman T Gerbode F and Selzer A Malfunction of mitral ball valve prosthesis due to thrombosis *Am J Cardiol* 29 95 1972
- 31 Hultgren H N and Hubis II A phonocardiographic study of patients with Starr Edwards mitral valve prosthesis *AM HEART J* 69:1306 1965
- 32 Craig E Hutchin P and Sutton R Impaired function of cloth covered Starr Edwards mitral valve prosthesis *Circulation* 41:441 1970
- 33 Leachman R D and Cocklinos D V P Absence of opening click in dehiscence of mitral valve prosthesis *N Engl J Med* 281 161 1969
- 34 Hylen J C Kloster F E Herr R H et al Sound spectrographic diagnosis of aortic ball variance *Circulation* 39 849 1969
- 35 Lansing A M Unusual radiological sign of loose mitral valve prosthesis *Radiology* 88:189 1967
- 36 Barold S S Javier R P and Linhart J W Factitious mitral insufficiency *N Engl J Med* 279 810 1968
- 37 Garamella J J Lynch M F Schmidt W R et al Fatal clotting of the Starr Edwards mitral ball valve nineteen months postoperatively *J Thorac Cardiovasc Surg* 47 613 1964
- 38 McHenry M M Smeloff E A Fong W Y et al Critical obstruction of prosthetic heart valves due to lipid absorption by silastic *J Thorac Cardiovasc Surg* 59:413 1970
- 39 Spencer F C Trinkle S K and Reeves J T Successful replacement of a thrombosed mitral ball valve prosthesis *JAMA* 194 191 1964
- 40 Carey J S and Huges R A A complication of mitral valve replacement with caged lens prosthesis *Ann Thorac Surg* 6:77 1968

Hemopericardium tamponade and effusion Cardiac tamponade may occur in association with any cardiac injury which produces hemopericardium and repeated pericardiocentesis may be required.²² Tamponade may appear as long as a week or more following injury and prompt pericardiocentesis may be required to avert catastrophe. Although pericardiocentesis is widely acclaimed in the treatment of localized penetrating wounds to the heart its role in the treatment of most myocardial wounds due to blunt trauma is limited by the usually greater extent of myocardial damage present and by the nonlocalized nature of the damage. On occasion retained blood can lead to late occurring chronic pericarditis with constriction months or years afterwards which may necessitate pericardiectomy.

Pericarditis Some degree of traumatic pericarditis usually from pericardial and/or myocardial ecchymosis or hematoma is found at postmortem in most cardiac injuries due to blunt chest trauma. Although occasionally reported to be of clinical significance resolution from this disorder is the rule. Should however intra-cardiac surgery be required later obliteration of the pericardial sac may cause technical difficulties at operation especially if the cardiac defect has resulted in cardiac enlargement.¹⁹

Delayed recurring pericardial effusions acute pericarditis and chronic constrictive pericarditis may follow both closed and penetrating cardiac injuries.²¹ Among five cases reported by Guest and colleagues²³ three followed blunt injury. Why some patients go on to develop recurrent pericarditis or constriction is not well understood but Tabatznik and Isaacs²⁴ believed recurrent inflammation to be present secondary to an altered autoimmune process which sometimes responded to corticosteroid administration. The vast majority of patients with late-occurring constrictive pericarditis however show no demonstrable cause. Extensive calcification of the pericardium²⁴ has been described as an additional long term complication in this setting and Mannix and Dennis²⁵ have recommended pericardiectomy in cases which become symptomatic.

Myocardial contusion In the context of

modern cardiac surgery myocardial contusions may seem trivial and of little clinical consequence yet well documented instances of death following this lesion have been recorded for hundreds of years.²⁶ The first case of unquestionable myocardial contusion reported by Akenstide¹⁰ in 1764 was in a 14 year-old boy who died 6 months after injury. However it was not until 1935 that Bright and Beck¹⁸ working with a trauma preparation in dogs first called attention to the significant pathophysiologic consequences that form the basis of our modern concepts concerning this lesion.

To understand the signs and symptoms of myocardial contusion it is helpful to review the pathological changes in the myocardium. The external appearance of the heart may show little if any evidence of injury yet a small area of subepicardial hemorrhage may overlay a much larger area of intramural contusion. Extensive contusion of the intraventricular septum can exist in the absence of other gross evidence of injury. The transition from normal to damaged myocardial tissue is usually abrupt in contusion in contradistinction to myocardial infarction where more gradual transitions occur. Otherwise the histological changes are similar in the two lesions with myocardial cell necrosis infiltration of polymorphonuclear leukocytes absorption of hemorrhage and healing by scar formation occurring in both. Ultimate healing of both contusions and infarcts may be virtually identical and differentiation between the two processes at a later time must rely on history condition of the coronary arteries and other evidence.²⁷ When extensive damage is present throughout the myocardium early death is the rule although some later deaths have been recorded. Causes of death include ventricular fibrillation¹⁹ prolonged cardiac standstill²⁸ myocardial rupture and/or heart failure.²⁹

Clinical manifestations and diagnostic criteria for myocardial contusion follow closely from the similarity with myocardial infarction. In a group of verified cases gathered from the literature fully 70 per cent of patients with myocardial contusion reported chest pain identical with that customarily associated with the onset of coronary occlusion.³¹ Most patients had concomitant chest wall injuries with resultant musculo

Table I *Classifications of cardiac lesions in nonpenetrating chest trauma*

| |
|---|
| I Pericardium |
| A Disruption |
| B Hemopericardium and tamponade |
| C Pericarditis |
| II Myocardium |
| A Contusion |
| B Rupture |
| C Septal perforation |
| D Late aneurysm |
| III Valves chordae tendinae and papillary muscles |
| IV Coronary arteries |
| A Contusion and thrombosis |
| B Laceration |

usual, yet occasionally such accidents were recognized^{10, 11} and lethal arrhythmias and myocardial rupture were described.^{4, 12, 13} With the advent of the automobile however high speed deceleration impact injuries have markedly increased and currently represent by far the commonest cause of this type of lesion. Vehicular impact abrupt transfers of kinetic forces to the victim and sudden decelerations of bodily viscera and blood column momentum all operate sequentially to produce cardiac damage. The extent and severity of this injury is determined by several biomechanical interactions including the magnitude of deceleration, the total duration of exposure to the forces of deceleration, and the rate of time change of deceleration, according to Newtonian laws of dynamics

$$F = MA$$

or in the case of deceleration $F = MD$

$$\text{and } D = V_2 - V_1/t$$

where F is the resultant of forces acting on the viscera, M is the scalar mass of the viscera involved, D is the magnitude of deceleration, V_2 and V_1 are terminal and initial velocities respectively, and t is the time duration of velocity change.

Stapp¹⁷ and Lasky and colleagues¹⁸ have also described an additional hydraulic ram effect as contributing to the development of heart damage in selected injuries in which abdominal and lower extremity compression may upwardly displace abdominal contents and produce cardiac injury in the absence of actual thoracic trauma. Bright and colleagues¹⁹ Aren-

berg²⁰ and others^{21, 22} observed internal thoracic damage to be less in cases of nonpenetrating chest trauma in which skeletal rib fractures have occurred suggesting a protective influence of the thoracic wall in dissipating energy impact forces. This is supported by the data of Dunseth and Ferguson²³ who noted in cases of severe myocardial damage—i.e., 17 cases of ventricular septal rupture—that only 3 had sternal or rib fractures.

Classification of cardiac injuries

The lesions to be discussed are categorized by type and location in Table I. While some structures appear more resistant to damage than others, injuries to all structures have been reported both singly and in combination. When multiple the physiological consequences of each injury may interact to either preserve function or cause deterioration and/or death especially where pericardial involvement exists.

Pericardial disruption. How often pericardial rupture occurs in survivors is unknown. Parmley and colleagues⁴ found pericardial disruption in 71 of 346 autopsy cases. A higher incidence of pericardial tears was reported by the present authors⁴ in an experimental animal model preparation in 14 of 18 dogs receiving sublethal blunt chest trauma. Almost all rents were transverse and extended across the upper base of the heart near the reflection of the visceral and parietal pericardium. The clinical implications and sequelae of this rupture as an isolated lesion are the potential danger of cardiac herniation²⁴ with catastrophic hemodynamic compromise and possible death.

Boyd and Strieder²⁵ pointed out that when myocardial and pericardial defects coexist particularly with bleeding survival was largely dependent upon the relative size of the pericardial tear or conversely upon the degree to which the integrity of the pericardial sac was preserved to prevent exsanguination. With large tears death was inevitable. With sufficiently small disruptions however adequate tamponade may control myocardial bleeding long enough to permit surgical intervention. If both pericardial and myocardial wounds are small stabilization without treatment may proceed to complete recovery.

presence of cyanosis heart failure dyspnea or orthopnea Interestingly coronary vaso dilators and anticoagulants have proved of little benefit for this lesion

Myocardial rupture Myocardial rupture is the lesion most frequently found at autopsy following fatalities due to non penetrating chest trauma In addition to violent impact injuries closed chest mas sage in cardiopulmonary resuscitation has added to the frequency of this lesion In order of frequency rupture of the right ventricle occurs most often followed by the left ventricle right atrium and left atrium⁴ Its occurrence in relation to nonpenetrating chest trauma carries a much graver prognosis than with penetrating wounds primarily due to the greater extent of damage Penetrating defects of the myocardium have been frequently repaired successfully by surgical correction whereas closed chest lesions have rarely been corrected One series reported only 30 of 132 patients with ventricular chamber rupture to survive 30 minutes or longer following trauma¹⁹ Recently however encouraging reports have appeared documenting survival in a small series of surgically repaired atrial ruptures²⁰⁻²² In these patients hemopericardium tamponade and cardiac failure were tolerated for longer periods of time thus allowing for more optimal surgical interventions including cardiopulmonary assist devices

Physical findings of myocardial rupture in those patients surviving long enough to reach the hospital are characteristic of cardiac tamponade Depressed arterial blood pressure distended neck veins with an elevated central venous pressure muffled heart sounds pericardial friction rub and an enlarged cardiac shadow on chest film are all expected findings The electrocardiogram shows diffuse low voltages

Treatment consists of emergency surgery Too few survivors have been encountered to establish the benefit of interim pericardiocentesis but this seems doubtful as a therapeutic tool since blood loss is too massive Time and facilities permitting extracorporeal assist devices seem of great potential merit but in view of the fact that 75 per cent of high speed accident injuries occur in or near communities with popula-

tions of 2 500 people or less⁴ undue assistance upon having these sophisticated devices present before attempting repair may result in further loss of life

Surgical techniques found to be of use in penetrating wounds of the heart should also prove effective with nonpenetrating cardiac injuries The widest possible exposure is mandatory Digital occlusion of ventricular perforations and occlusive clamps applied to atrial defects will control bleeding while sutures are applied In inaccessible areas the insertion of a balloon catheter through the tear with inflation and traction may be helpful in controlling bleeding during repair

Septal perforation Despite its seemingly protected anatomical position the inter ventricular septum appears acutely susceptible to injury secondary to nonpenetrating chest trauma²³⁻²⁵ Predilection for septal perforation particularly near the apex is high²⁶ Many reports of this lesion are described in the literature^{10 27 28 29} Inkley and Barry²⁴ noted in animal studies an increased risk of vulnerability when the traumatic blow was interjected late in diastole or early systole²⁴

Diagnosis may be difficult since rupture of the chordae tendinae and/or papillary muscles with mitral or tricuspid incompetence severe myocardial contusion with congestive heart failure or pre existing congenital defects all may mimic acquired septal perforation An easily palpable precordial thrill and early holosystolic murmur maximal in the third and fourth intercostal spaces at the left sternal border are the characteristic findings but may be absent in the presence of shock The electrocardiogram may show left axis deviation left anterior hemiblock and/or right bundle branch block indicating multifascicular injury to conduction pathways Currents of injury³⁰ and several rhythm disorders^{2 30 31} have also been described Cardiac catheterization affords the only reliable means of establishing the correct diagnosis³²

Although small defects may close spontaneously the treatment of choice is surgical closure particularly in the presence of cardiac failure^{33 34} There have now been reported more than 20 surgical repairs for traumatic perforations of the interventricular septum^{45 35 36 37 100 101} Peirce and col

skeletal pain, but many of those with myocardial bruises had, in addition readily distinguished angina like symptoms which interestingly were not relieved by coronary vasodilating drugs

Diagnosis of myocardial contusion is rarely obvious on physical examination alone in the absence of major complications such as pericardial tamponade^{28 46 48} cardiac arrhythmias^{49 52} or heart failure^{44 45}. Tachycardia is invariably present in this setting but may be in response to associated injuries as well as heart damage

Twelve lead electrocardiography has proved to be the most reliable means of making the diagnosis of myocardial contusion in patients with thoracic injury. Despite possible errors in specificity^{53 54} electrocardiography offers an accurate practical and dependable means of evaluating the injured patient. The time course of changes in the electrocardiogram following cardiac trauma is not completely known but from the data available in the absence of arrhythmias fully a third of patients in the first 24 to 48 hours after injury will develop ST segment and T wave changes similar to myocardial ischemia and/or infarction^{57 58 59}. The duration of these changes varies but may persist for as long as a month following injury. These observations have been duplicated in several animal studies^{45 47 60 61} in which similar electrocardiographic changes were reported including transient elevations of the ST segment and T waves suggesting a current of injury pattern

Electrocardiographic changes which persist for periods of longer than a month and become permanent suggest lesions more complicated than simple contusion and include extensive myocardial scarring with possible left ventricular aneurysms and coronary artery injury which will be discussed in later sections

A wide range of rhythm disorders and conduction disturbances have also been documented, both in clinical and in animal studies. Cardiac arrhythmias have included premature ventricular contractions (the commonest type of dysrhythmia) complete cardiac standstill, second third and varying degrees of heart block, atrioventricular dissociation with or without interference,

ventricular tachycardia and fibrillation and atrial tachycardia flutter, and fibrillation^{19 24 42 49 51 62 67}. Conduction disturbances have included right and left bundle branch blocks and interventricular conduction delays^{52 68 71}. Aside from a weak correlation between these disorders and severity of injury, their mechanism of development is not understood despite extensive autopsy analyses of clinical and animal material^{11 44}. Direct response to injury interplay of neurohumoral reflexes coronary spasm and effects of intramyocardial hemorrhage have all been suggested^{19 42 65 72 74}

Other laboratory studies have not proved of much help in the diagnosis of myocardial contusion. Serum enzyme changes are frequently elevated in concomitant injuries of the liver lung bone brain and skeletal muscle but specificity for cardiac injury is essentially masked^{37 59 75}. Shubin and Weil¹⁶ and Vessel and associates⁷⁷ showed significant elevations of serum glutamic oxaloacetic transaminase (SGOT) and lactic dehydrogenase (LDH) in hemorrhagic shock per se which would limit their usefulness should major blood loss develop in association with cardiac injury. Studies in combat casualties⁷⁸ showed elevations in SGOT and LDH not only in shock but also in metabolic acidosis and lactic acidemia, which would further lessen their usefulness in the diagnosis of myocardial contusion

Other laboratory tests such as leukocyte count and sedimentation rate also lack sufficient specificity for myocardial contusion but may be helpful in following the clinical course of the patient to determine optimal times at which physical activity may be resumed. From a medicolegal standpoint all laboratory aids which can be employed with little risk to the patient should be used if the clinician believes they may help in assessing residual damage to the myocardium

In general the treatment of myocardial contusion is similar to that of myocardial infarction. Bed rest is most important usually for a 2 to 4 week period followed by a program of graded ambulation. Specific drug therapy should be strictly supportive. Digitalis is only effective in the control of heart failure or supraventricular tachyarrhythmias. Oxygen is indicated in the

results¹²¹ and prosthetic valve replacement is now recommended as the procedure of choice¹²²

Injuries of the mitral valve may involve tears of the anterior and/or posterior valvular leaflets but more commonly cause ruptures of the chordae tendinae and papillary muscles^{123, 124, 125} The mitral valve appears most vulnerable to rupture during diastole when the heart is distended. Rupture of a mitral valve chordae tendinae or papillary muscle is accompanied by the development of precordial pain, breathlessness, palpitation, shock or fulminating left heart failure. A precordial thrill with a loud harsh holosystolic murmur maximal at the apex and radiating to the axilla is present on physical examination. An apical diastolic murmur indicating torrential return of blood across the mitral valve may also be heard and cardiac enlargement usually develops with time. Rupture of a papillary muscle is catastrophic and leads to rapid development of left heart failure with death in hours or days. Early operation affords the best chance of survival. Methods of repair have historically been individualized with successes reported both with valvuloplasty and prosthetic valve replacement^{126, 127} The latter is favored currently by most clinicians.

Traumatic injury of the tricuspid valve with dysfunction and insufficiency is fortunately rare. In 1848 Todd¹²⁸ first reported this injury and accurately described the pathophysiology leading to its damage. He concluded that violent external cardiac compression plus sudden pulmonary outflow obstruction put great strain on the chordae tendinae and papillary muscle attachments to the tricuspid valve leading to tears and rupture. Jahnke and associates¹²⁹ as recently as 1967 reviewed the world literature on traumatically acquired tricuspid insufficiency and found 17 cases. Since then Morgan and Foraker¹³⁰ have reported on three others and Vladoff and Desforges¹³¹ have reported on one. The majority of patients were men; most trauma involved automobile accidents and the described lesions included ruptured papillary muscles, chordae disruption, or detachment, valve leaflet rupture and complete valve destruction.

Symptoms of tricuspid incompetence

vary from mild to severe and include fatigue, massive peripheral congestion with ascites and edema and occasionally dyspnea and orthopnea. Similar to mitral valve dysfunction, disruption and tearing of the papillary muscles following injury has produced the most fulminant tricuspid insufficiency with rapid clinical decline. Physical findings include central venous hypertension with obvious V wave pulsations in the neck, hepatic pulsations with positive hepatojugular reflex and a loud holosystolic murmur heard along the lower left sternal border which increases with inspiration.

In the group reviewed by Jahnke and colleagues¹²⁹ 10 had electrocardiograms and 8 showed complete or incomplete right bundle branch block. On fluoroscopic examination, marked paradoxical pulsation of the right atrium and superior and inferior vena cavae were seen. Cardiac catheterization done in 9 demonstrated ventricularized atrial pressures in systole with giant V waves ranging from 8 to 28 mm Hg. Six patients were operated on between three months and ten years after injury and three received prosthetic valves. Primary repair of the papillary muscles has also been attempted in a few,^{132, 133, 134} but results have proved poor and valve replacement in sufficiently compromised patients now appears to be the procedure of choice.^{135, 136, 137, 138, 139}

Coronary arteries The question of whether nonpenetrating chest trauma can cause direct injury to the coronary arteries either by thrombosis, laceration or occlusion has been strongly debated in the literature^{140, 141, 142} for some time. Evidence from several sources would support such a relationship^{143, 144} but further and more direct documentation, particularly with coronary arteriography, is required. Nevertheless, when electrocardiographic evidence of myocardial infarction is obtained shortly after injury in a young individual with no prior history of heart disease and these changes persist permanently, the differential diagnosis should include traumatically acquired coronary artery hemorrhage and/or occlusion.

At the present time, too few numbers of unequivocal cases representing this syndrome are currently available to recommend definitive courses of therapy. How

leagues,⁴ who reviewed the histories of 18 patients not operated on found that 10 died of congestive heart failure within 15 days of their injuries. This experience serves to emphasize the urgency of proceeding promptly with cardiac catheterization and surgical repair.

Late aneurysm. This complication of non-penetrating injuries to the heart, if accompanied by advanced symptoms of failure, has rarely been treated successfully to date. Its true incidence is unknown since many of those which cause little disability probably go undiagnosed. In those instances where myocardial aneurysm is recognized it is not certain whether myocardial contusion directly¹⁰² or in combination with coronary artery disease acquired either secondarily as a response to trauma or developed separately on an atherogenic basis has caused formation of the aneurysm.

In addition to true aneurysm myocardial contusion and small ruptures may also produce pseudoaneurysm. O'Reilly¹⁰³ reviewed 15 cases of ventricular aneurysm following nonpenetrating chest trauma, nine of which were in children and found that where the character of the lesions was clearly described three were true and five were false or pseudoaneurysmal.

Morbidity and mortality statistics in both types of aneurysm are high with fatal complications including myocardial rupture and tamponade, cardiac failure, systemic emboli and arrhythmias causing sudden death.¹⁰⁴ Lyons and Perkins¹⁰⁵ reported the first operative treatment for post-traumatic left ventricular aneurysm in 1958, and preliminary results from studies reported by Killen and associates¹⁰⁶, Aronstam and colleagues,¹⁰⁴ and Pupello and co-workers¹⁰⁷ as well as from the increasing experience with management of post-infarction aneurysm¹⁰⁸ suggest this mode of therapy may soon become the accepted treatment of choice.

Valves, chordae tendinae and papillary muscles. Prior to the advent of open heart surgery, injuries to cardiac valves and their supporting structures were irreparable, and except in instances where the hemodynamic sequelae were mild, therapy was ineffective. Some form of corrective treatment is now available for almost all such injuries and early diagnosis may avoid

serious disability or death. Untreated rupture of a valve is usually followed by rapid progression of congestive heart failure with eventual demise within a year or two.

Historically, these lesions have generated considerable curiosity and interest in the literature beginning with Barrie,¹⁰⁹ who in 1881 demonstrated in cadavers that extreme pressures were required to rupture cardiac valves. Dufour¹¹⁰ as early as 1891, experimentally produced aortic rupture in dogs. A variety of traumatic mechanisms postulated to cause these lesions were investigated including direct chest wall impact, abdominal and lower extremity compression, various strains and blast pressure from explosions.^{111, 112} Kulbs¹¹³ concluded that only modest increases in intra-aortic pressures were required to cause aortic valve rupture and ascribed the greater frequency of left-sided valve injuries to higher pressures in these chambers. Pre-existing condition of the valves is also important as Freidberg¹¹⁴ reported a greater incidence of traumatic valve ruptures in those patients with pre-existing disease in either the mitral and/or aortic valves.

In order of frequency traumatic impact injuries produce rupture of the aortic valve most often followed next by damage to the mitral valve.^{115, 116} Howard¹¹¹ has reviewed the literature on aortic insufficiency and has found 44 believed to be due to trauma or strain. Of these 22 valves showed rupture along the base at the annular attachment of the cusps, 11 had tears in one or more of the cusps, in 2 of which all 3 cusps were torn and 10 of the valves were frankly detached from the aorta. This complication occurred overwhelmingly in men.

In aortic lesions there is a typical high-pitched diastolic blowing murmur heard with the usual circulatory hemodynamic changes associated with aortic regurgitation.¹²⁰ A distinctive musical murmur described as 'sea gull' or 'cooing dove' in character has also been attributed to aortic valve rupture but is not invariably present. Cardiac catheterization and supraventricular angiography are essential in identifying this lesion and early recognition is mandatory in order to avert the complications of massive cardiac insufficiency with left ventricular failure. Primary repair of the damaged valve was attempted early with poor

- Engl J Med 260 1139 1961
- 32 Guest J L Hall D P Vela T J and Ellison R C Late manifestations of trauma to the pericardium Surg Gynecol Obstet 120 187 1965
- 33 Tabatznik B and Isaacs J P Postpericardiotomy syndrome following traumatic hemopericardium Am J Cardiol 7 83 1961
- 34 Warburg E Traumatic armour heart AM HEART J 49 633 1955
- 35 Mannix E P and Dennis C A A study of the surgical treatment of chronic pericardial effusion and cardiac tamponade J Thorac Surg 29 381 1955
- 36 Kinsane R W Traumatic heart disease non penetrating injuries Circulation 6 421 1952
- 37 DeMuth W E and Zinsler H F Myocardial contusion Arch Intern Med 116 434 1965
- 38 DeMuth W E Baue A E and Odom J A Contusions of the heart J Trauma 7 443 1967
- 39 Kahn M H and Kahn S Cardiovascular lesions following injury to the chest Ann Intern Med 2 1013 1929
- 40 Leinhardt H D Acute coronary thrombosis in industry direct nonpenetrating injuries with report of cases Arch Intern Med 70:33 1942
- 41 Moritz A R Pathology of trauma Philadelphia 1947 Lea & Febiger Publishers pp 143 149
- 42 Kujala F and Strauss L H Heart and trauma further experimental research Klin Wochenschr 11 1572 1932
- 43 Lounimo I Heart injury after blunt thoracic trauma an experimental study on rabbits Acta Chir Scand Suppl 380 1 1968
- 44 Anderson R H Nonpenetrating injuries of the heart Br Med J 2 307 1940
- 45 Pearce H C Dabbs H and Rawson F Isolated rupture of the ventricular septum due to nonpenetrating trauma Arch Surg 77 87 1958
- 46 Shackelford R T Hydropericardium report of a case with summary of the literature JAMA 96 187 1931
- 47 Rajasingham A S Massive hemopericardium with recovery after paracentesis Br Heart J 1:181 1939
- 48 London R F and London B B The electrocardiographic signs of acute hemopericardium Circulation 25:780 1962
- 49 Kinsane R W Fidler R S and Koons R A Electrocardiographic changes following external chest injury to dogs Ann. Intern. Med 11 907 1937
- 50 Sgier L H Traumatic injury of the heart AM HEART J 30 459 1945
- 51 Taylor H M Transient cardiac arrhythmia induced by nonpenetrating trauma to the chest AM HEART J 46 557 1953
- 52 Gozo E G Cohen H C and Pick A Traumatic bifascicular intraventricular block Chest 61:294 1972
- 53 Reid J M and Baird W L M Crushed chest injury some physiological disturbances and their correction, Br Med J 1 1105 1965
- 54 Dolara A Morando I and Pampaloni M Electrocardiographic findings in 98 consecutive nonpenetrating chest injuries Dis Chest 32:30 1967
- 55 Berry F B Chest injuries Surg Gynecol Obstet 70 413 1940
- 56 Barber H Electrocardiographic changes due to trauma Br Heart J 4 83 1943
- 57 Barber H The effects of trauma direct and indirect on the heart Q J Med (n.s.) 13:137 1944
- 58 Lepeschkin M Modern electrocardiography vol 1 Baltimore 1951 The Williams & Wilkins Company
- 59 Watson J H and Bartholomae W M Cardiac injury due to nonpenetrating chest trauma Ann Intern Med 52 871 1960
- 60 Randles F S Gorham L W and Dresbach M Changes in the RST component of the electrocardiogram produced by experimental rupture of the auricle of the dog's heart and by pericardial injection AM HEART J 9 333 1934
- 61 Rosenkranz K A Meier G and Humperdinck H Electrocardiogram and serum enzyme activity following experimental traumatic heart injuries in relation to pathologic-anatomic findings Monatsschr Unfallheilkd 68:337 1965
- 62 Rosenow W Heart block in a child of 10 years following trauma to the precordium Am J Dis Child 28 594 1924
- 63 Warburg E Traumatic heart lesions London 1938 Oxford University Press pp 1676-1868
- 64 Coffin T H Rush H P and Miller R F Traumatic complete heart block of 18 years duration with review of literature Northwest Med 40 195 1941
- 65 Scherf D Blumenfeld S and Yildiz M Experimental study on intracardiac tachycardia and cardiac trauma Cardiologia 40 37 1964
- 66 Sims B A and Geddes J S Traumatic heart block Br Heart J 31 140 1969
- 67 Bharati S Chervony A Gruhn J Rosen K M and Lev M Atrial arrhythmias related to trauma to sinoatrial node Chest 61 331 1972
- 68 Sweeney P J Traumatic bundle branch block Clin J 7 218 1948
- 69 Rankin T J and Patterson J W Transient intraventricular block in cardiac contusion AM HEART J 43 103 1952
- 70 Hale H W and Marlin J W Myocardial contusion Am J Surg 93 558 1957
- 71 Zinsler H F and Thund G S Right bundle branch block after nonpenetrating injury to the chest wall JAMA 207 1913 1969
- 72 Schlomka G and Hinrichs A Experimental proceedings on the influence of blunt chest trauma on the electrocardiogram Z Gesamte Exp Med 81 43 1932
- 73 Schlomka G and Schmitz M Experimental proceedings on the influence of blunt chest trauma on the electrocardiogram II Z Gesamte Exp Med 83:179 1932
- 74 Scherf D and Boyd L J Cardiac changes due to nonpenetrating trauma in cardiovascular diseases their diagnosis and treatment

ever treatment should be supportive and similar to the management of other forms of acute myocardial infarction. Whether future management for this complication will include the newer surgical revascularization procedures such as aortocoronary saphenous vein bypass graftings remains speculative at this time.

Summary

Nonpenetrating chest trauma with injury to the heart has become increasingly common, particularly as a result of deceleration injuries in modern, high speed vehicular accidents. The present report reviews the currently available information concerning the several varieties of cardiac lesions which are incurred, including damage to the pericardium, myocardium, valvular structures, and coronary arteries. Although highly lethal if neglected and often obscured by associated trauma, cardiac damage can generally be estimated quickly with a proper index of suspicion and relatively simple diagnostic tools. Modern medical and surgical techniques in the appropriate clinical settings have proved curative.

REFERENCES

- 1 Committee on Trauma and Committee on Shock. Accidental death and disability. The neglected disease of modern society. Washington D C 1965. National Academy of Sciences p 5
- 2 Spelman J. Personal communication 1966
- 3 Committee on Trauma and Committee on Shock. Accidental death and disability. The neglected disease of modern society. Washington D C 1966. National Academy of Sciences p 21
- 4 Parmley FF, Manion WC and Mattingly TW. Nonpenetrating traumatic injury of the heart. *Circulation* 18:375 1958
- 5 Silbernagel WM and Fidler RS. Intracardiac traumatic lesion of the heart. *Am HEART J* 26:129 1943
- 6 Hunt WE. Spontaneous rupture of the heart in a newborn infant. *Arch Dis Child* 27:291 1952
- 7 McInray RA and Graham ALM. Rupture of the fetal heart during labor. *Arch Dis Child* 28:201 1953
- 8 Price AC, Van Praagh R, Sears WP and Nadas AS. Post traumatic left ventricular myocardial infarction and rupture in infancy. *J Pediatr* 72:656 1968
- 9 Beck CS. Contusions of the heart. *JAMA* 104:109 1935
- 10 Akenside M. Account of blow upon heart and its effects. *Phil Trans R Soc. London* 1:64 p 353
- 11 Hewitt P. Rupture of the heart and large vessels: the result of injuries. *London Medical Gazette* 4 (ns) 8:0 1847
- 12 Fischer G. Die Wunden des Herzens und des Herzbeutels. *Langenbeck Arch Klin Chir* 9:571 1868
- 13 Kellert E. Traumatic rupture of the heart: report of a case with uninjured chest wall. *J Lab Clin Med* 2:726 1916-17
- 14 Kissane RW, Koons RA and Fidler RS. Ruptured aortic valve following explosion. *Am HEART J* 12:31 1936
- 15 Stein W and Revitch E. Traumatic rupture of the right ventricle. *Am HEART J* 21:103 1942
- 16 Miller G III and Rueb AE. Blunt thoracic trauma producing heart laceration: case report. *Ann Surg* 166:852 1967
- 17 Stapp JP. Gravitational stress in aerospace medicine. Boston 1961. Little, Brown & Company pp 168-188
- 18 Lasky I I, Nahum AM and Siegel AW. Cardiac injuries incurred by drivers in automobile accidents. *J Forensic Sci* 14:13 1969
- 19 Bright EF and Beck CS. Nonpenetrating wounds of the heart: Clinical and experimental studies. *Am HEART J* 10:293 1935
- 20 Arenberg H. Traumatic heart disease: clinical study of 250 cases of nonpenetrating chest injuries and their relation to cardiac disability. *Ann Intern Med* 19:326 1943
- 21 Ferré GA and Steward WD. Cardiac contusion. *Chn Orthopedics and Related Research* 53:123 1967
- 22 Rotman M, Peter RH, Sealy WC and Morris JJ. Traumatic ventricular septal defect secondary to nonpenetrating chest trauma. *Am J Med* 48:127 1970
- 23 Dunseth W and Ferguson T II. Acquired cardiac septal defect due to thoracic trauma. *J Trauma* 5:112 1965
- 24 DeMuth WF, Lerner EH and Liedtke AJ. Nonpenetrating injury of the heart: an experimental model. *J Trauma* (In press)
- 25 Hoffman K T. Traumatic rupture of the pericardium with heart luxation. *Thorax chirurgie* 14:67 1966
- 26 Boyd TF and Stedner JW. Immediate surgery for traumatic heart disease. *J Thorac Cardiovasc Surg* 50:3 1965
- 27 Goodland MJ, Bloomer WE and Goodyer AV. Recurrent pericardial effusion after nonpenetrating chest trauma. *N Engl J Med* 263:874 1960
- 28 Tygart RL, Mitchell JA and Glas WW. Cardiac contusion complicated by cardiac tamponade. *J Mich State Med Soc* 61:313 1962
- 29 Hatcher CM and Bahnsen HT. Cardiac contusion: puncture and tamponade. *Am J Surg* 105:458 1963
- 30 Therkelsen F. Surgical repair of traumatic ventricular septal defects. *Acta Chir Scand* 119:372 1960
- 31 Pastor RH and Betts RH. Late symptoms due to traumatic heart contusion.

- 111 Howard C P Aortic insufficiency due to rupture by strain of a normal aortic valve *Can Med Assoc J* 19:12 1928
- 112 Frothingham C and Haas G M Rupture of normal chordae tendinae of the mitral valve *AM HEART J* 9 492 1934
- 113 Kujbs F Experimentelle untersuchungen ueber herz und trauma *Mitt Grenzgeb Med Chir Jena* 19 678 1909
- 114 Freiberg C K Diseases of the heart Philadelphia 1969 W B Saunders Company p 1697
- 115 Bushong H B Traumatic rupture of aortic valve report of two cases one proved and another probable example of this condition *Ann Intern Med* 26:125 1947
- 116 Leonard J J Harvey W P and Hufnagel C A Rupture of the aortic valve a therapeutic approach *N Engl J Med* 2:2 208 1935
- 117 Probst W L and McCormack L J Rupture of the aortic valve *Circulation* 13:750 1956
- 118 Gregersen H Traumatic rupture of the aortic valve report of a case *Acta Chir Scand* 129 669 1964
- 119 Najafi H Dye W S Javid H Hunter J A and Julian O C Rupture of an other wise normal aortic valve report of two cases and review of the literature *J Thorac Cardiovasc Surg* 57 1968
- 120 Levine H J Roberts W C and Morrow A G Traumatic aortic regurgitation *J Cardiol* 10 732 1962
- 121 Spurny O M and Hara M Rupture of the aortic valve due to strain *Am J Cardiol* 8:125 1961
- 122 Beall A C and Shirkey A L Successful surgical correction of traumatic aortic valve regurgitation *JAMA* 187 507 1964
- 123 Glendy R F and White P D Nonpenetrating wound of heart Rupture of papillary muscle and confusion of heart resulting from external violence case report *AM HEART J* 11:366 1936
- 124 Menges H Ankeney J L and Hellerstein H K Ruptured mitral chordae tendinae *Circulation* 30 8 1964
- 125 McLaughlin J M Cowley N A Smith G and Matheson N A Mitral valve disease from blunt trauma *J Thorac Cardiovasc Surg* 48 261 1965
- 126 Bouvrain Y Four cases of rupture of the chordae tendinae of the mitral valve treated by valvuloplasty *Bull Soc Med Hop Paris* 116 567 1965
- 127 Sanders C A Scannell J G Harthorne J W and Austen W G Severe mitral regurgitation secondary to ruptured chordae tendinae *Circulation* 31 306 1965
- 128 Szekacs A Traumatic rupture of papillary muscle with unrecognized cardiac tamponade *J Forensic Sci* 11 174 1966
- 129 Todd R B A case of rupture of the chordae tendinae of the tricuspid valve of the heart with remarks *Dublin Q J Med Sci* 5 1 1848
- 130 Jahnke E J Nelson W P Ashby G V and Fitzgibbon G M Tricuspid insufficiency The result of nonpenetrating cardiac trauma *Arch Surg* 9:840 1967
- 131 Morgan J R and Forker A P Isolated tricuspid insufficiency *Circulation* 43:559 1971
- 132 Vladoff I M and Desforges G Cardiac injuries due to nonpenetrating thoracic trauma *Ann Thorac Surg* 14 504 1977
- 133 Cooley D A Henley W S Amadi H II and Chapman D W Ventricular aneurysm following myocardial infarction *Ann Surg* 150 395 1959
- 134 Osborn J R Jones R C and Jahnke E J Traumatic tricuspid insufficiency Hemodynamic data and surgical treatment *Circulation* 30:217 1964
- 135 Bjork V O Traumatic rupture of the tricuspid valves *Thoraxchirurgie* 12 368 1965
- 136 Brandenburg R O McGoon D C Campeau L and Guiliani E R Traumatic rupture of the chordae tendinae of the tricuspid valve *Am J Cardiol* 11 911 1966
- 137 Shabetai R Adolph R J and Spencer F C Successful replacement of the tricuspid valve 10 years after traumatic incompetence *Am J Cardiol* 18 916 1966
- 138 Salzer J Weintraub R Lower R and Eldridge F Isolated tricuspid insufficiency report of a case with valve replacement *Am J Cardiol* 18:921 1966
- 139 Shean Ming L Sako Y and Alexander C S Traumatic tricuspid insufficiency *Am J Cardiol* 26 200 1970
- 140 Boas E P Angina pectoris and cardiac infarction from trauma or unusual effort—with consideration of certain medicolegal aspects *JAMA* 112 1887 1939
- 141 Levy H Traumatic coronary thrombosis with myocardial infarction—postmortem study *Arch Intern Med* 11 261 1949
- 142 Lehman H J Sundquist A H and Giddings L W Coronary thrombosis with myocardial infarction secondary to nonpenetrating injury of the chest wall *AM HEART J* 47 470 1954
- 143 Montz A R Trauma stress and coronary thrombosis *JAMA* 156 1306 1954
- 144 Levy R L de la Chapelle C and Richards D W Heart disease in drivers of public motor vehicles as a cause of highway accidents *JAMA* 184 481 1963
- 145 Campbell M Angina pectoris following a crushing accident *Br Heart J* 1 177 1939
- 146 Warburg E Myocardial and pericardial lesions due to nonpenetrating injury *Br Heart J* 2:171 1940
- 147 MacDonald D Repeated and fatal coronary thrombosis in a young man *JAMA* 116:2846 1941
- 148 Harthorne J W Kantrowitz P A Dismore W H and Sanders C A Traumatic myocardial infarction Report of a case with normal coronary angiogram *Ann Intern Med* 66:341 1967
- 149 Stewart J S S Primary traumatic coronary thrombosis *Br Med J* 1:1739 1967
- 150 Jones F L Jr Transmural myocardial necrosis after nonpenetrating cardiac trauma *Am J Cardiol* 26:419 1970

- London 1948 Medical Books Ltd Publishers pp 273-276
- 75 Wroblewski F The clinical significance of transaminase activities of serum *Am J Med* 27:911 1959
 - 76 Shubin H and Weil M H Acute elevation of serum transaminase and lactic dehydrogenase during circulatory shock *Am J Cardiol* 11:327 1963
 - 77 Vessell E S Feldman M P and Frank E D Plasma lactic dehydrogenase activity in experimental hemorrhagic shock *Proc Soc Exp Biol Med* 101:644 1959
 - 78 Sleeman H K Simmons R L and Heisterkamp C A Serum enzymes in combat casualties *Arch Surg* 98:1272 1969
 - 79 Desforges G Ridder W P and Ienoci R J Successful suture of ruptured myocardium after nonpenetrating injury *N Engl J Med* 252:1567 1955
 - 80 Bogedarm W Carpathios J Suu D V and Moots M F Traumatic rupture of myocardium Successful surgical repair *JAMA* 197:154 1966
 - 81 Borja A R and Lansing A M Traumatic rupture of the heart a case successfully treated *Ann Surg* 171:438 1970
 - 82 Siderys H and Strange P S Rupture of the heart due to blunt trauma *J Thorac Cardiovasc Surg* 62:184 1971
 - 83 Pollock B E Markelz R A and Shuey H E Isolated traumatic rupture of interventricular septum due to blunt force *Am Heart J* 43:273 1952
 - 84 Cary F H Hurst J W and Arentzen W R Acquired interventricular septal defect secondary to trauma report of 4 cases *N Engl J Med* 258:1355 1958
 - 85 Rubenstein P and Levinson D S Acquired interventricular septal defects due to myocardial infarction and nonpenetrating trauma to the chest *Am J Cardiol* 7:277 1961
 - 86 Miller D R Crockett J C and Potter C A Traumatic interventricular septal defect—review and report of two cases *Ann Surg* 155:172 1962
 - 87 Kanber G J Fort M L Treger A Meadows W R and Sharp J T Left ventricular-right atrial canal with aortic incompetence of probable traumatic origin *Am J Cardiol* 20:879 1967
 - 88 Guilfoil P H and Doyle J T Traumatic cardiac septal defect report of case in which diagnosis is established by cardiac catheterization *J Thorac Surg* 25 510 1953
 - 89 Paulin C and Rubin I L Complete heart block with perforated interventricular septum following contusion of the chest *Am Heart J* 52 940 1956
 - 90 Stern W R and Stoddard L D Traumatic interventricular septal defect of heart *Am Heart J* 63 821 1962
 - 91 Tascon A M and Rostrup O Rupture of the interventricular septum due to blunt trauma *Can J Surg* 8:197 1965
 - 92 Williams G D Hara M and Bulloch E Traumatic ventricular septal defect *Am J Cardiol* 18 907 1966
 - 93 Stinson E H Rowles D F and Shumway N E Repair of right ventricular aneurysm and ventricular septal defect caused by nonpenetrating cardiac trauma *Surgery* 64 1022 1968
 - 94 Inkley S R and Barry F M Traumatic rupture of interventricular septum proved by cardiac catheterization *Circulation* 11 916 1958
 - 95 Dolara A La comunicazione interventricolare da rottura isolata del setto nei traumi contusivi del cuore *Minerva Medicoleg* 84:110 1964
 - 96 Meister S G and Helfant R H Bedside diagnosis of ruptured ventricular septum with mitral insufficiency *N Engl J Med* 287:1024 1972
 - 97 Cleland W P Ellman P Goodman J and Hallman A Repair of ventricular septal defect following indirect trauma *Br J Dis Chest* 55:117 1961
 - 98 Desforges G and Abelmann W H Interventricular septal defect due to blunt trauma report of a case repaired surgically under total cardiopulmonary bypass *N Engl J Med* 268 128 1963
 - 99 Turney S Z Mathai J Singleton R and Cowley R A Traumatic ventricular septal defect surgical repair in two patients *Ann Thorac Surg* 13:136 1972
 - 100 Gahagan T and Green E W Repair of complicated defect in cardiac septum after nonpenetrating trauma *JAMA* 194 301 1965
 - 101 Green L Oakley C M Davies D M and Cleland W P Successful repair of left ventricular aneurysm and ventricular septal defect after indirect injury *Lancet* 2 984 1965
 - 102 Joachim H and May A T A case of cardiac aneurysm probably of traumatic origin *Am Heart J* 2 682 1927
 - 103 O'Reilly R J Gregorio K and Spellberg R D Traumatic pseudoaneurysm of left ventricle *Am J Dis Child* 120:252 1970
 - 104 Aronstam M Strader L D Geiger J P and Gomez A C Traumatic left ventricular aneurysms *J Thorac Cardiovasc Surg* 59 239 1970
 - 105 Lyons C and Perkins R Resection of a left ventricular aneurysm secondary to cardiac stab wound *Ann Surg* 147:256 1958
 - 106 Kullen D A Gobbell W G France R and Via V A Post traumatic aneurysm of the left ventricle *Circulation* 39:101 1969
 - 107 Pupello D F Daily P O Stinson E B and Shumway N E Successful repair of left ventricular aneurysm due to trauma *JAMA* 211 826 1970
 - 108 Cooley D A and Hallman G L Surgical treatment of left ventricular aneurysm Experience in excision of post infarction lesion in 80 patients *Progr Cardiovasc Dis* 9:722 1968
 - 109 Barre E Recherches cliniques et experimentales sur les ruptures valvulaires du coeur *Rev Med Paris* 1:132 1881
 - 110 Dufoeur C Des insuffisances aortiques d'origine traumatique *T* 1896 7

V J me 86
V mb 5

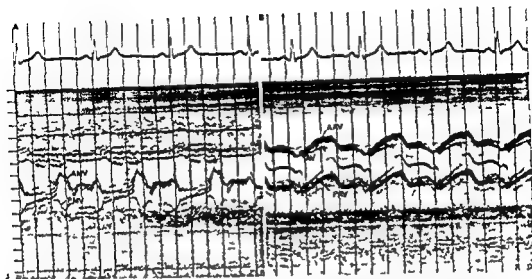


Fig 1 A and B A Mitral and B aortic echograms from a normal subject. The individual echoes from the valve leaflets are sharp and free of any thickening or shaggy echoes. Valvular motion is unrestricted. AMV = anterior mitral valve leaflet. PMV = posterior mitral valve leaflet. AAV = anterior aortic valve leaflet. PAV = posterior aortic valve leaflet.

surgery or their hearts were examined at autopsy. All patients had numerous blood cultures.

Results

Four of the eight patients were found to have positive blood cultures. *Streptococcus viridans* was the organism in three of the patients. The blood culture in the fourth patient grew enterococcus. The four patients with negative blood cultures had received prior antibiotic treatment before transfer to this facility but had clinical evidence of bacterial endocarditis and were appropriately treated. At autopsy or surgery five patients had vegetations on the aortic valve and three had vegetations on the mitral valve. Of the three patients with mitral valve lesions the vegetations were on the anterior leaflet in one, the posterior leaflet in another and on both leaflets on the third. The vegetations in all of the patients varied from 2 to 8 mm in size.

Fig 1 shows mitral valve and aortic valve echograms from a normal subject. Figure 1A shows the anterior and posterior mitral valve leaflets while the anterior and posterior aortic valve leaflets are demonstrated in Fig 1B. It should be noted that these echoes are sharp and free of any thickening or shaggy echoes attached to the valve. Fig 2 is a composite of several

views of the aortic valve echogram from a patient in whom the aortic valve was involved with vegetations. The echograms are characterized by a non uniform thickening in several areas of the valve. In many areas the thickened valve has a shaggy appearance. Although these shaggy echoes are seen in each view of the aortic valve the individual echograms are slightly different demonstrating the non uniformity of this abnormality. These different echograms also illustrate why it is usually necessary to scan the entire valve to be certain to visualize any abnormality. In Fig 2C the aortic valve appears to be free of any abnormal echoes during systole and the abnormal shaggy material occurs during diastole. In Fig 2B the abnormal echoes seem to be seen best during systole. A consistent observation in all of the views is that despite the apparent thickening of the aortic valve leaflets the valve motion remains normal.^{2,3} The other four patients with pathologically proved vegetations on the aortic valve had echocardiograms similar to those exhibited in Fig 2.

Fig 3 is a mitral valve echogram from a patient who has valvular vegetations of the anterior leaflet of the mitral valve. There is uneven thickening noted under the anterior leaflet. This thickening can be noted both during systole and diastole. Despite this

Fundamentals of clinical cardiology

Echocardiographic manifestations of valvular vegetations

James C Dillon, M D
Harvey Feigenbaum M D
Lee L Konecke, M D
Richard H Davis, M D
Sonia Chang A B
Indianapolis, Ind

The presence of vegetations on cardiac valves is a hallmark of bacterial endocarditis.¹ This study was undertaken to see if echocardiography could visualize such vegetations and if so to determine what size these vegetations would have to be in order to be seen by echocardiography. Hopefully if these vegetations could be visualized by echocardiography this information might not only be useful diagnostically, but it might provide some information concerning the pathogenesis and natural history of this disease.

Method

Eight patients with either surgically or autopsy proved valvular vegetations were evaluated using echocardiography. The echocardiograms were obtained using an Ekoline Mark IIA echocardiograph coupled to an Electronics for Medicine recorder A 2.25 MHz, 0.5 inch transducer, focused at 7.5 cm was employed in all patients. The patients were examined in the recumbent position. The transducer was

placed in the third or fourth intercostal space along the left sternal border. The transducer was directed posteriorly and slightly medially in order to pick up the mitral valve echoes. The transducer then was rotated superiorly and medially towards the right shoulder to pick up the aortic root and valve. If the aortic valve cusps were not obtained in this position the transducer was usually moved up one interspace and again pointed superiorly and medially. Both the mitral and aortic valve echoes were obtained in all eight patients. In doing the echocardiographic examination emphasis was placed on an M mode scanning technique whereby the direction of the transducer is changed while recording continuously.² This technique permits the recording of many views of each valve in a continuous recording. It avoids the problem of having the echogram vary with the direction of the transducer and also provides a more complete examination of the particular valve in question.

All patients either underwent open heart

From the Department of Medicine, Indiana University School of Medicine and the Krannert Institute of Cardiology, Marion County General Hospital, Indianapolis, Ind.
Supported in part by the Herman C. Krannert Fund, United States Public Health Service Grants PHS-HE-09815-07, HE-6308, HTS-5363, HE-5749 and the Indiana Heart Association.
Received for publication Oct. 23, 1972.
Reprint requests to Dr. James C. Dillon, 110 Fessler Hall, 1100 W. Michigan St., Indiana University School of Medicine, Indianapolis, Ind. 46202.

November 1973 Vol. - p. 698

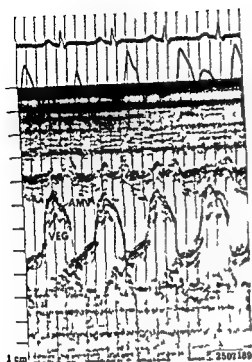


Fig 3 Mitral valve echograms from a patient with vegetations on the anterior mitral leaflet. There are multiple echoes (VEG) on the anterior leaflet (AMV) originating from the vegetations. Despite this apparent thickening of the leaflet the valve motion remains unrestricted and there is actually some coarse fluttering of the valve during diastole.

a shaggy appearance and is non uniform with respect to the areas of the valve which are involved.

Whether or not these echocardiographic findings are specific for valvular vegetations remains to be proved. There does not seem to be any problem in confusing these echograms with the usual causes of thickened mitral or aortic valves.^{2,9} Fig 6 shows the mitral valve echograms from two patients with mitral stenosis. In one patient there is minor thickening of the leaflets with abnormal motion and a decreased diastolic slope. In Fig 6B the mitral valve is heavily calcified with a large band of echoes noted throughout the valve. The dense echoes are not spotty and the valve motion is definitely abnormal.^{4,7}

Fig 7 contains echocardiograms from a patient who has another condition that could be confused with valvular vegetations. This patient has a left atrial myxoma. There are several differentiating features

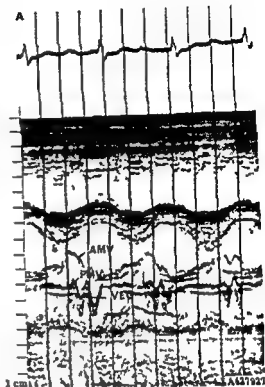


Fig 4 Echocardiogram from a patient with vegetations involving the posterior mitral leaflet. The vegetations are recorded as a block of shaggy echoes (VEG) attached to a freely movable posterior mitral valve leaflet (PMV). The anterior mitral valve leaflet (AMV) is thin and apparently uninvolved with vegetations.



Fig 5 Surgical specimen of the mitral valve from the patient whose echogram is seen in Fig 4. The large vegetation involving the posterior leaflet is easily seen (arrow). Note also the perforated anterior leaflet and the ruptured chordae.

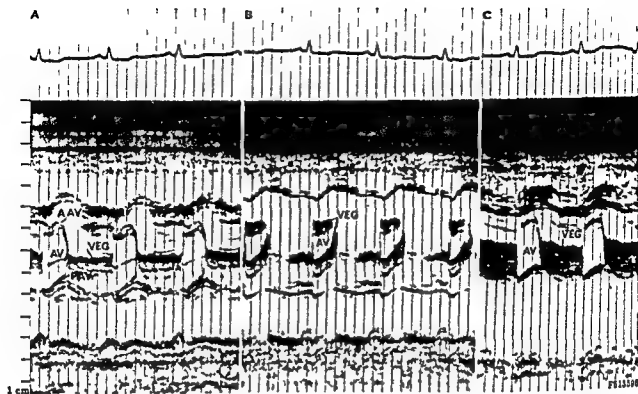


Fig 2 A through C Three aortic valve echograms from a patient with vegetations on the aortic valve. The location and appearance of the vegetations vary depending upon the angle of the transducer and how the ultrasonic beam strikes the valve. Despite the apparent valvular thickening the motion of the leaflets remains unrestricted. AAV = anterior aortic valve leaflet AV = aortic valve PAV = posterior aortic valve leaflet VEG = vegetation

apparent increase in the amount of valvular echoes, the motion of the valve remains unrestricted and even exhibits some coarse fluttering during diastole.⁴⁷ Fig 4 is another echocardiogram from a patient with bacterial endocarditis and vegetations on the mitral valve. In this patient, the vegetations were confined solely to the posterior leaflet. The large shaggy echo attached to the echo of the posterior leaflet is easily seen (VEG). Again the motion of the posterior mitral leaflet is good⁴⁸ despite the presence of this large echo producing mass. Fig 5 is a photograph of the mitral valve following surgical removal from the patient whose echogram is seen in Fig 4. The large vegetation on the posterior leaflet is noted by the arrow. The echogram from the patient who had vegetations on both mitral leaflets demonstrated shaggy echoes on both leaflets with the motion of the leaflets remaining unrestricted. In all eight patients, the location of the echocardiographic valvular abnormalities correlated with the location of the vegetations at surgery or autopsy.

We were able to obtain serial echograms on two of the eight patients with vegetations. In these two patients, the intensity and size of the echoes presumably originating from the valvular vegetations, increased with time despite the patients' apparent bacteriologic cure both clinically and by culture of the surgical specimen.

In one patient who proved to have bacterial endocarditis confined to the aortic valve, the echocardiogram was considered to be positive for vegetations before the diagnosis was expected clinically. This patient had chronic renal failure and was undergoing echocardiography to rule out pericardial effusion.

Discussion

The results of this study seem to indicate that echocardiography can detect valvular vegetations provided that they are at least 2 mm in diameter. The echocardiographic manifestations of these vegetations seem to be an apparent thickening of the valve with no restriction in motion of the leaflets. In addition, the thickening frequently has

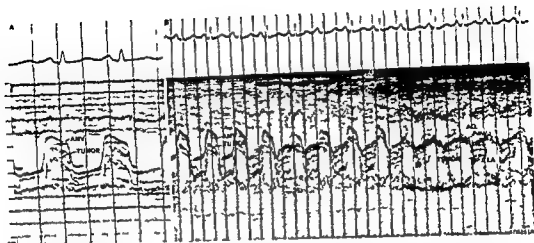


Fig 7 A and B Echocardiograms from a patient with a left atrial tumor. The tumor echoes in echogram A superficially resemble echoes arising from vegetations. One differentiating feature is that the tumor obscures the echo from the posterior mitral leaflet. In echogram B directing the ultrasonic beam toward the left atrium (LA) one continues to record tumor echoes in the left atrium indicating that the echoes most likely originate from a mass in the left atrium rather than from vegetations attached to the valve leaflet.

echocardiography to be done serially provides an excellent opportunity to study the natural history of patients with treated bacterial endocarditis. It is quite possible that many patients with apparently cured bacterial endocarditis go on to form thickened and significantly deformed valves.

Unfortunately it could not be determined from this study as to when in the course of illness the vegetations appeared because the exact onset of the illness was not known in many of the patients. We have examined some patients with clinically evident bacterial endocarditis in whom we were unable to demonstrate any valvular vegetations. None of these patients went to surgery or autopsy. This observation raises the question as to how much time is required from the onset of the illness until vegetations can be seen echocardiographically. In addition it is possible that in the successfully treated case of endocarditis the vegetations may never get large enough to be recorded on the echogram. These questions obviously must be answered before the echocardiographic technique can be fully evaluated as to its usefulness in the diagnosis of bacterial endocarditis. Thus although this study demonstrates that echocardiography can detect valvular vegetations secondary to bacterial endocarditis and that such

examinations may prove to be useful in enhancing our understanding as to the role of vegetations in this disease, the specificity of this echocardiographic finding must yet be proved and the exact role of this technique in the diagnosis of bacterial endocarditis must be studied further.

Summary

Eight patients with autopsy or surgically proved valvular vegetations were examined using echocardiography. Five of these patients had lesions on the aortic valve and three had lesions on the mitral valve. The echocardiographic finding in these patients was a non uniform thickening of valve leaflets which exhibited unrestricted motion. Often the abnormal echoes which produced the thickened valve had a shaggy appearance. In all eight patients the location of the echocardiographic abnormality correlated with the anatomic findings at surgery or autopsy. In one patient the diagnosis of bacterial endocarditis was first suspected following the echocardiographic examination and only subsequently was a heart murmur heard. These findings indicate that echocardiography may play a useful role in elucidating the pathological anatomy of the bacterial endocarditis with vegetation, however the length of time from the onset of clinical

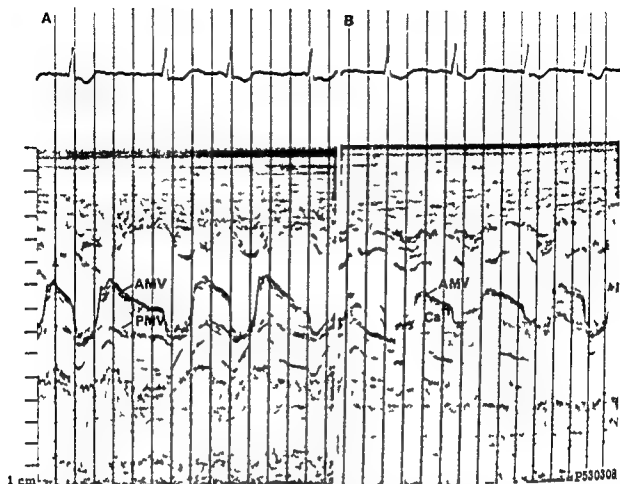


Fig 6 A and B Mitral valve echograms from two patients with mitral stenosis. The valve in echogram A is mildly fibrotic whereas the mitral valve in B is heavily calcified. Both valves exhibit uniform thickening of the leaflets and the calcium (Ca^{++}) in B has a shaggy appearance. Both valves can be distinguished from patients whose thickened valves are due to vegetations by the abnormal motion characteristic of mitral stenosis.

which distinguish this echogram from those taken of patients with valvular vegetations.^{10,11} In Fig 7A the posterior mitral valve leaflet is not seen because it is obscured by the tumor. In addition the extra echoes do not seem to be directly attached to the valve leaflets. Possibly the most convincing evidence that the patient in Fig 7 had a left atrial tumor and not valvular vegetations is the fact that the tumor continues to be seen as the ultrasonic beam is directed towards the left atrium. Valvular vegetations are attached to the valve and have not been noted in the left atrium.

We have had one patient with myxomatous degeneration of the mitral valve who exhibited thickening of the posterior mitral valve on the echocardiogram. The leaflet motion continued to be good and the valve did not look like one which was scarred by rheumatic disease. The appearance of

the valve was different from those with vegetations in that the thickening was fairly uniform and the echoes did not appear shaggy. However it should be noted that not all of the patients with vegetations had the shaggy echoes on the valve (Fig 3). Thus there could conceivably be some confusion between the echograms of patients with vegetation and those with myxomatous degeneration and thickened valves.

The fact that two patients had a progressive increase in the intensity and size of the echoes originating from the vegetations despite an apparent bacteriologic cure raises several interesting questions. The development of increased echoes could represent a normal healing process and not necessarily a progression of active infection. Obviously, further studies are necessary to answer this question but the ability of

Complications of aortocoronary artery bypass surgery

Alfred J Kaltman MD
New York NY

The effectiveness of aortocoronary artery bypass grafting in the treatment of occlusive coronary disease is related to selectivity of patients, surgical technique and skills and the subsequent course of the disease process. Expertise and experience with heart lung bypass operations in general have minimized operative risks but there are specific complications of coronary artery grafting which must be considered in the evaluation of these procedures.

Myocardial infarction in the intraoperative and early and late postoperative periods, fate of the arteries proximal and distal to the site of anastomosis, continued patency of the grafts and the progression of disease in grafted and ungrafted vessels are important factors in the late morbidity and death of the patient.

Myocardial infarction

Variability in interpretation of early postoperative electrocardiograms has made the frequency of acute myocardial infarcts during surgery difficult to determine. Persistence of the serum isoenzymes of lactic dehydrogenase and the cardiac specific myoglobin fractions of creatine phosphokinase may supplement the diagnosis of equivocal electrocardiographic changes.

The operative mortality rate of most reported series varied from less than 3 per cent to about 10 per cent. Evidence of acute myocardial infarction as manifest by the appearance of abnormal Q waves was present in more than 50 per cent of the deaths in whom early postoperative electrocardiograms were available. Anatomic evidence of myocardial infarction was found in almost 60 per cent at autopsy. In some instances of necropsy proved necrosis the electrocardiograms were not definitely diagnostic.

Approximately 15 per cent of patients surviving coronary bypass grafting may have definite electrocardiographic evidence of acute myocardial infarction. If occurrence of left bundle branch block and left anterior hemiblock are included, 17 to 18 per cent of survivors have myocardial necrosis during surgery.

The total frequency of acute myocardial infarction during coronary bypass grafting appears to be in the range of 20 per cent of all patients, both survivors and those succumbing to surgery. Since repolarization abnormalities of the electrocardiogram are almost universal in the early postoperative period, subendocardial and small intramural infarctions may be missed by the strict criteria of Q wave appearance and

From the Department of Medicine, New York University Medical Center, New York, N.Y.

Received for publication July 11, 1973.

Reprint requests: Alfred J. Kaltman, MD, Department of Medicine, New York University Medical Center, 550 First Avenue, New York, N.Y. 10016.

illness to echocardiographic diagnosis remains unknown

REFERENCES

- 1 Freidberg C K Disease of the heart 3rd ed Philadelphia and London 1966 W B Saunders Company ■ 1382
- 2 Feigenbaum H Clinical application of echocardiography Progr Cardiovasc Dis 14:531 1972
- 3 Gramiak R and Shah P M Echocardiography of the normal and diseased aortic valve Radiology 96:1 1970
- 4 Edler I Herz C H Gustafson A et al The movements of the heart valves recorded by means of ultrasound Nord Medic 64:1178 1960
- 5 Joyner C R Jr Reid J M and Bond J P Reflected ultrasound in the assessment of mitral valve disease Circulation 27:503 1963
- 6 Segal H I Likoff W and Kingsley H Echocardiography clinical application in mitral stenosis JAMA 195:161 1966
- 7 Zaky A Nasser W K and Feigenbaum H Study of mitral valve action recording by reflected ultrasound and its application in the diagnosis of mitral stenosis Circulation 37 789 1968
- 8 Dillon J C Haine C L Chang S and Feigenbaum H Use of echocardiography in patients with prolapsed mitral valve Circulation 43:503 1971
- 9 Duchak J M Chang S and Feigenbaum H The posterior mitral valve echo and the echocardiographic diagnosis of mitral stenosis Am J Cardiol 29 628 1972
- 10 Schattenberg T T Echocardiographic diagnosis of left atrial myxoma Mayo Clin Proc 43 620 1968
- 11 Wolfe S B Popp R L and Feigenbaum H Diagnosis of atrial tumors by ultrasound Circulation 39 615 1969
- 12 Nasser W K Davis H H Dillon J C Tavel M E Helmen C H Feigenbaum H and Fisch C Atrial myxoma phonocardiographic echocardiographic hemodynamic and angiographic features in nine cases AM HEART J 83 810 1972

internal mammary artery more closely approximates the caliber of the distal coronary artery than the larger saphenous vein has furthered the recent interest in this technic. In addition to the internal mammary direct anastomosis to the left anterior descending and circumflex arteries free grafts of this artery have been used for aorto right coronary artery bypass. Indeed in some instances the splenic artery has been brought through the diaphragm to the right coronary artery. Too few arterial grafts of these various types have been done to assess adequately their effectiveness and complications.

Progression of coronary artery disease after bypass grafts

Progression of stenosis in the coronary artery proximal to graft anastomosis apparently is quite common. This portion of the vessel may become totally occluded in a large number of cases. It has been demonstrated in as many as 60 per cent of instances in which vein grafts were non patent within two weeks of operation and in 20 per cent of those patients whose grafts closed during the first year. Indeed proximal coronary arteries had progressed to occlusion in 30 per cent of one series of patients with grafts patent for one year.

The presence of an apparently functioning graft does not preclude the progression of stenosis in the section of the artery distal to the anastomosis. This has been reported in 8 per cent of one surgical group. In that same series 13 per cent of patients revealed progression of atherosclerosis in non grafted vessels after one year.

Obviously the underlying disease process atherosclerosis is not basically affected by these surgical interventions. On the other hand alteration of the flow and hemodynamics of the vasculature by interposition of grafts has an effect on the stresses and responses of the affected coronary arteries. A minimal existing process conceivably can be accelerated and earlier stenosis result from a combination of thrombotic and atherogenic factors. This may even be augmented by increasing flow through the distal vessel beyond a critical point. The high incidence of proximal arterial occlusion suggests that a preferential flow through the graft may

cause a steal syndrome with deprivation of flow subsequent thrombosis and occlusion of these diseased arteries. As yet there are no meaningful data regarding the mechanisms of these observations and further study is necessary.

Conclusions

At the present time the operative mortality rate of aortocoronary artery bypass grafting ranges from less than 3 per cent to about 10 per cent. More than half of these deaths are due to myocardial infarction. Another 17 to 18 per cent of surviving patients may develop myocardial infarction during the year following operation.

About 20 per cent of saphenous vein grafts fail to remain patent for one year. Closure usually is associated with return of angina untoward change in clinical status and possible myocardial infarction. Many physiological pathological and technical factors are relevant to graft patency. Of prime importance is the establishment of blood flow of more than 40 ml per minute through the grafts and adequate size of the recipient artery at the site of anastomosis.

Medial hypertrophy and subintimal fibrous hyperplasia of vein grafts, the experimental greater durability of arterial grafts and the reported higher long term patency of internal mammary artery to left anterior descending coronary artery bypasses have stimulated current interest in the use of arteries rather than veins for grafting material. In addition to direct internal mammary anastomoses to the left anterior descending coronary free arterial grafts and direct splenic artery bypasses have been utilized for left circumflex and right coronary artery bypass procedures.

Progression of the underlying disease process atherosclerosis is not halted by these surgical techniques. Total occlusion of the coronary artery proximal to the site of graft insertion is extremely common regardless of whether the graft remains open or closed during the first postoperative year. The recipient coronary artery distal to the anastomosis may stenose in a smaller number of patients about 8 per cent in one group restudied. Atherosclerosis had progressed in non grafted coronaries after one year in 13 per cent. The change

the number of undiagnosed areas of necrosis may be underestimated

The incidence of operative myocardial infarction is significantly related to the severity of anginal symptoms, number of grafts inserted, and prolongation of heart lung bypass pump time beyond two hours. Statistically significant differences were not apparent in regard to smoking, previous myocardial infarction, or abnormal lipoprotein profile.

During the year following surgery late myocardial infarction unrelated to graft closure is difficult to document. Occasionally silent myocardial infarction can occur in the presence of patent grafts and a few patients may remain relatively asymptomatic despite occlusion of one or even all of the grafts. Usually graft closure is associated with dysrhythmias, return of angina, hypotension, a distinct deterioration in clinical status, and clear cut new electrocardiographic abnormalities consistent with acute myocardial necrosis.

Patency of grafts

Clinical response of angina correlates well with graft patency. Very few patients remain completely asymptomatic in the presence of closed grafts. Early closure has been associated with lack of symptomatic improvement and occasionally with impairment of function. Recurring angina after initial relief, sudden arrhythmias and transient hypotension can occur during the first postoperative year and may require early restudy to determine the adequacy of the revascularization procedure. Reoperation following clinical failure of bypass grafts may be feasible and necessary.

Loss of patency of inserted grafts remains a significant complication. In all series there is both an early and late closure rate which approaches 20 to 25 per cent within the first year. Many physiological, pathological and technical factors have been implicated. In general patency has been attributed to those grafts with favorable blood flow of more than 40 ml per minute established at surgery and adequate caliber of the recipient artery with good distal runoff.

There is an inverse relation of flow to length and the cross sectional area of the

conduit created. Short, small caliber grafts have more rapid flow rates. An advantage of increased velocity of flow is to reduce the tendency for platelet and fibrin deposition and decrease thrombus formation.

Wall tension of the grafts is proportional to their length and to the size of the lumen. Greater tension is developed as the radius is increased and the grafts are elongated. In addition to the effects of enhanced tension in regard to possible development of medial hypertrophy of the grafts in the future, increased resistance may be a factor in thrombus formation within the grafts.

Coronary flow occurs predominantly in diastole. During systole, myocardial tissue pressure around small distal coronary arteries may reduce transmural pressure and temporarily occlude them, contributing to thrombus in traumatized grafts or inefficient anastomoses. Competitive flow between the proximal artery and a graft or even between different grafts and the presence of collateral circulation may jeopardize graft patency.

Although atherosclerosis has not been recognized, subintimal fibrous hyperplasia has developed in saphenous vein grafts. Reaction to stress of arterial pressure on the vein wall and a possible water hammer effect created by distal stenosis may be responsible. Other factors are ischemia of the vein and mechanical injury at the time of harvesting.

Problems with maintenance of function in vein grafts have focused attention on the use of internal mammary artery anastomoses to the left anterior descending coronary artery. A 97 per cent long term patency rate has been claimed. However, many of these patients required additional vein grafts to left circumflex and/or the right coronary arteries, only 70 per cent of which were found functional at the time of restudy. Furthermore, operating time is longer with its attendant higher complication rate.

The internal mammary artery has a relatively small diameter with an inherent high velocity, an aid to patency. On the other hand, it may not always provide adequate flow for distal arteries. The fact that arterial grafts experimentally have been more durable than vein grafts—that the

Annotations

Whither electrocardiography?

The art or science whichever one prefers of electrocardiography is now some 60 years old. Many advances have occurred since the pioneering studies of Einthoven and yet today electrocardiography still has many limitations. Reasons for this less than satisfactory state are such things as variations in body habitus from patient to patient, the placement of precordial electrodes and the response of the recording instrument—to name a few.

The position of the precordial electrodes has been standardized by the British and American Heart Associations but in actual practice these positions are seldom accurately ascertained and certainly the positions are not reproduced accurately by technicians in the recording of daily electrocardiograms. *It would obviously help to mark the positions which were actually used by a dye or by silver nitrate if multiple electrocardiograms are expected to be taken.* In lieu of this a different positioning of the electrodes over the left precordium to encompass an area instead of a line would be of help. This might include a high V_4 and V_5 lead with a V_6 lead in the center like the five spot on a die.

Let us consider the recording instrument. The original string galvanometer as devised by Einthoven was a most satisfactory instrument provided the electrodes were properly applied to achieve low skin resistance. The response was linear from DC to several hundred cycles per second. However the instruments tended to be bulky, difficult for technicians to use and the strings were frequently broken. The records required photographic processing as well. To get around some of these problems direct writing instruments using vacuum tube amplifiers and heated stylus recording on heat sensitive paper were introduced some 30 years ago. Instruments of this type have been improved by the use of solid state components and response electrical safety, time decay and similar functions are better. The fact remains however that salesmen, technicians and even physicians can easily change the response of the instrument to provide cleaner records by increasing the pressure of the recording stylus on the paper. Thus the instrument no longer meets the specifications set by the manufacturer and supposedly met when the instrument was delivered to the consumer.

In 1967 a special Committee of the American Heart Association¹ proposed minimal standards for electrocardiographic instruments. This received some attention from the manufacturers and now a

few companies produce instruments which meet some if not all of these minimal standards.

Accuracy is of prime importance in the frequency range from DC (flat response) to perhaps 2 000 Hz but there is no universal agreement on these figures particularly the upper limit. (AHA minimum standards call for a flat response from 0.14 cps to 50 Hz with a 3 db falloff at 100 Hz.) The higher the frequency response the more problems arise from 60 Hz interference and from muscle tremor and most studies in the high frequency range above 130 cps have been limited to research investigations.

There are however certain limiting factors in the direct writing electrocardiograph which make it less than an ideal instrument for accurate recording of the electrocardiogram. These include the mass and inertia of the stylus and the friction of the stylus on the paper. There will always be certain compromises which must be made between cost, size, accuracy, ease of operation, patient safety, sensitivity to outside interference and the like.

The low frequency response of the instrument (0 to 5 Hz) in view of today's knowledge is probably of equal if not greater importance. Errors in the 0 to 5 Hz band are common due to paper friction or stiction and poor amplifier response to minute signals in this range. This is especially important when one considers that probably 90 per cent of abnormalities noted in the electrocardiogram are confined to this range: the P wave, the PR interval, the S-T segment, the T wave and the U wave. It is especially important to recall that repolarization is beginning in some cells even before the end of depolarization (QRS) and much information may be present in the S-T segment if it is accurately recorded.

This brings us to the considerations of what may be done to improve the quality of direct writing electrocardiograms without unduly affecting cost and other factors mentioned previously.

One way to improve low frequency accuracy by lessening the effect of paper stiction and stylus friction and amplifier distortions in the ≈ 0.1 mv range is to amplify the voltage and record at 2 to 20 times the normal standardization. It is obvious that this will usually throw the QRS complexes off the paper which may damage the stylus or activate a protect circuit. It is however possible using special circuitry to clip the QRS complexes and get accurate recording of the low frequency components (Fig. 1). Accuracy in timing can be achieved

in hemodynamics and flow patterns within the coronary circulation provoked by the interposition of bypass grafts is complex and requires further study in order to understand any direct relationship to possible acceleration or deceleration of the basic occlusive arterial disease.

These complications of aortocoronary artery bypass surgery are important determinants of postoperative morbidity and death. They will influence the subsequent course of the patient and can change an excellent initial clinical response into an untimely death. Better understanding of the mechanisms of these complications undoubtedly will help reduce their incidence in the future. At the present time recognition of these events is necessary to afford the patient the best possible therapeutic management during the early and subsequent postoperative periods.

SELECTED REFERENCES

- Adam M, Mitchell B F, Lambert C J and Geisler G F Long term results with aorta to coronary artery bypass vein grafts. *Ann Thorac Surg* 14:1 1972
- Brewer D L, Bilbro R H and Bartel A G Myocardial infarction as a complication of coronary bypass surgery. *Circulation* 47:59 1973
- Hultgren H, Miyagawa M, Buck W and Angell M Ischemic myocardial injury during coronary artery surgery. *AM HEART J* 82:624 1971
- Urschel H C, Razzuk M A, Wood R E and Paulson D L Factors influencing patency of aortocoronary artery saphenous vein grafts. *Surgery* 72:1048 1972
- Groncin C M, Castonguay Y R, Lesperance J, Bourassa M G, Campeau L and Groncin P Attrition rate of aorta coronary artery saphenous vein grafts after one year. *Ann Thorac Surg* 14:223 1972
- Fleming R J, Johnson W D, Lepley D Jr, Tector A J, Walker J, Gale H, Beddingfield G and Manley J C Late results of saphenous vein bypass grafting for myocardial revascularization. *Ann Thorac Surg* 14:232 1972
- Green G E Internal mammary artery to coronary artery anastomosis. *Ann Thorac Surg* 14:260 1972
- Wlodaver Z and Edwards J E Pathologic changes in aortic coronary arterial saphenous vein grafts. *Circulation* 44:719 1971
- Aldridge H E and Trimble A S Progression of proximal coronary artery lesions to total occlusion after aortocoronary saphenous vein bypass grafting. *J Thorac Cardiovasc Surg* 62:7 1971
- Spencer F C, Green G E, Tice H A and Glassman E Bypass grafting for occlusive disease of the coronary arteries. *Ann Surg* 173:1029 1971

high frequency response with a flat response from DC to perhaps 2000 Hz Paper speed should be faster and accurately timed Amplitudes should probably be increased at least in those leads with very low voltage There should be a built in testing system which can check the continuity of all leads the function of networks used in unipolar lead current leakage frequency response and decay times

J Scott Butterworth MD
Ephraim Glassman MD
Department of Medicine
New York University School of Medicine
New York NY
Present address 104 E 40th St
New York NY 10016

Aspirin in the prevention of thrombosis

It has been demonstrated that aspirin inhibits the platelet release reaction which suggests that the prolongation of the bleeding time by aspirin is probably but not necessarily secondary to this action

This finding raised the possibility that aspirin might prove valuable in some pathological conditions involving thrombosis The idea needed exploring in view of the enormous medical social and economic importance of thrombotic conditions in the Western world and it was decided that the following trials should be carried out

Before their elected operation and daily for five days thereafter 303 patients were given 600 mg of aspirin or a placebo in a carefully controlled double-blind study carried out in four separate hospitals The patients legs were scanned daily after the injection of 125 I labelled fibrinogen for hot spots which indicate local fibrin deposition i.e. a thrombus The result was clear cut A dose of 600 mg of aspirin has no effect at all on the incidence of hot spots during the first five postoperative days and the incidence of clinically detectable thrombosis was too low for a meaningful comparison but the few cases observed occurred equally in the treated and placebo groups¹ Although platelets are believed to play a vital part in blood coagulation the extent to which they are involved in the deposition of isotopically labelled fibrinogen in the legs is of course not known However it can now be deduced with certainty that the release reaction and any other changes brought about by aspirin play no part in the formation of these thrombi after operations

This is a strong assertion particularly in the present climate of opinion about the role of platelets in clinically detectable thrombotic disease and the widespread use of aspirin clinically to protect patients from thrombosis So the following points must be made

Aspirin has no effect on the primary wave of platelet aggregation—stickiness—and red by adeno-

REFERENCES

- 1 Willems J L Pobleto P F and Pipberger H V Day to-day variation of the normal orthogonal electrocardiogram and vectorcardiogram *Circulation* 33 1057 1972
- 2 Kossmann C E Brady D A Burch G E et al Recommendations for standardization of leads and specifications for instruments in electrocardiography and vectorcardiography *Circulation* 33 583 1967

sine diphosphate so this mechanism could still be involved in venous thrombosis detected by the 125 I labelled fibrinogen method used in the study This can be resolved only by finding compound which interfere with this process and subjecting them to a similar clinical trial

Are the hot spots detected by the 125 I fibrinogen method a true reflection of clinical thrombosis? There is no doubt that the accumulation of fibrinogen in the leg veins does indicate a thrombus locally and this can usually be demonstrated by venography But there are certain important differences between thrombi detected by this method and the clinical disease we have known so long For example

hot spots occur frequently (in up to 71 per cent of some prolonged operations) Furthermore they are usually mild arise in the calf and are often found on the first postoperative day The vast majority are clinically silent and are associated with no known sequelae By contrast clinically detectable postoperative venous thrombi are usually recognized later are found in the thigh and of course occur much less frequently All the differences can be attributed to the fact that only a small proportion of 125 I fibrinogen type thrombi extend along the veins up into the thighs where they are found later and much less frequently So that although the very great value of the 125 I fibrinogen test in research and sometimes in clinical conditions is accepted it is pertinent to ask if it is too sensitive on the one hand or whether the condition we need to investigate is a succession of events which occur subsequent to the deposition of the original thrombus When one considers the most serious complication of postoperative thrombosis pulmonary embolism it is probably true that the 125 I fibrinogen test would be positive in all cases even in patients with silent pulmonary thrombi but this does not explain why it is positive in such very large numbers of patients who remain free from all symptoms

It might be suggested therefore bearing in mind

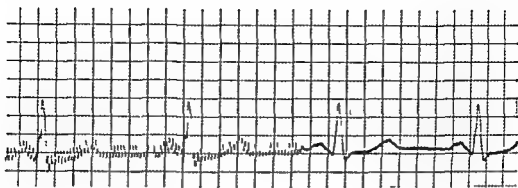


Fig 1 Example of an electrocardiograph lead recorded at 50 mm per second paper speed. A 60-cycle notch filter was inserted into the circuit after the first two complexes and the almost complete rejection of 60 Hz interference can be seen without significant alteration in the electrocardiographic complexes. The short black lines at the top of the tracing indicate 1-second time intervals. The paper speed here is approximately 51 mm per second—an error of 2 per cent. This figure can be used for corrections if necessary. This is more important at speeds of 100 mm per second as used in recording systolic time intervals.

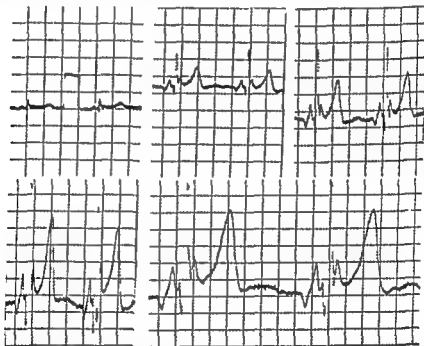


Fig 2 The top left box indicates an electrocardiographic lead at normal standardization of 1 millivolt equal 1 cm. The top middle box is the same lead amplified three times; the top right box is the same lead amplified six times; and the lower left box has an amplification of ten times. The above were recorded at a paper speed of 25 mm per second as indicated by the one second time intervals at the top of the tracings. The lower right box is the same as the lower left ($\times 10$) except that the paper speed is 50 mm per second. Many details in the P, QRS, and S-T segment can be seen which are not readily apparent in the standard lead.

by recording at paper speeds of 50, 75, or 100 mm per second provided the paper can be moved accurately. A time marker must be simultaneously recorded to assure this accuracy. More paper is used but it is only necessary to record a very few complexes at fast speeds and high amplifications.

Another way in which the records can be improved is by the use of sharp notch filters since the greatest interference by far encountered is 60 Hz. This will change the height of the QRS complexes to some extent, but it is doubtful if this will really limit the diagnostic importance (see Fig 2 with and without notch filter).

Other refinements consist in the recording of multiple simultaneous leads. This makes the interpretation of artifacts and arrhythmias much easier.

While there are other areas that could be mentioned, these points give an indication of the problems. As long as we accept the use of direct writing electrocardiography with the limitations inherent in it, it may be worthwhile to reconsider some of the concepts of standards such as electrode placement, paper speed, and amplification.

If it is really our intention to obtain the best possible recording of the electrocardiogram, then a suitable authoritative group should set standards of

connected peripheral portions depending upon the anterior subdivision, the posterior subdivision and a central radiation or plexus of ramifications coursing to the midseptal area.

These data were not entirely new. Several older papers reviewed by Rossi¹ reached similar conclusions. More recently Medrano and associates² observed the same distribution of the left bundle fibers in the dog. They also showed that it was possible to produce experimentally a concomitant block of both the anterior and posterior subdivisions without producing the classical pattern of left bundle branch block provided the midseptal ramification were not injured during the experimental procedure.³ Our results also fit in very well with the electrophysiological data obtained by Durrer and co-workers⁴ who found that the excitator of the endocardial surface of the left ventricle simultaneously starts at three widely separated areas which may reasonably be assumed to correspond to the termination zones of the three main portions of the left branch. In the light of these results the current concept which states that the left bundle branch is anatomically a bifascicular system appears tainted with over simplification.

Let us now turn toward the histopathological lesions underlying the patterns of hemiblocks. Recent observations indicate that it would be hazardous when one sees a tracing with LAHB or LPHB to draw hasty conclusions as to the precise topography of the partial bundle branch lesions responsible for the electrocardiographic abnormalities.⁵ In the above mentioned paper² we also reported on the histopathological features observed in ten patients with LAHB. In nine out of 10 instances conspicuous alterations of the left branch were depicted. They totally interrupted the anterior subdivision in five patients but were hardly ever limited to an anterior locus. The pathological process generally involved most of the left sided conduction system. Since this paper appeared several additional cases were studied. They brought supportive evidence in favor of our initial opinion which is shared by Rossi¹ and Blondeau and Lenegre.⁶ We may therefore conclude that the patterns of hemiblocks are manifestations of left bundle branch disease but that the underlying pathological lesions are generally much more widely distributed than expected from the electrocardiographic terminology of left anterior or posterior hemiblocks.

It is our belief that the problem of left intraventricular conduction disturbances should be reassessed keeping in mind these histological data. Indeed the validity of the term hemiblock which

etymologically implies that the bifascicular system of the left bundle branch system has been used as the unique basis for terminology should be questioned. Similarly further basic and clinical studies are still needed to determine the actual role played by the centro-septal fibers in the genesis of both the normal and pathological QRS.

The numerous paper that were devoted in recent years to partial left bundle branch blocks have opened a chapter which represents a most significant contribution to electrocardiology. This chapter should not be considered as completed. Further work remains to be done in this field.

J C Demoulin M D
Department of Morbid Anatomy
H E Kubitertus M D
Chercheur Qualifié du F.R.S.
Maître de Conférences
Division of Cardiology
Department of Medical Clinics and Semiology
University of Liège School of Medicine
Liège Belgium

REFERENCES

- 1 Pryor R and Blount M G. The clinical significance of true left axis deviation. *AM HEART J* 72:391 1966
- 2 Medrano G A, Brenes C F, De Michel A and Sodi Pallares D. El bloqueio simultáneo de las subdivisiones anterior y posterior de la rama izquierda del haz de His (bloqueo bifascicular) y su asociación con bloqueos de la rama derecha (bloqueo trifascicular). *Arch Int Cardiol Mex* 40:752 1970
- 3 Durrer D, van Dam R Th, Freud G E, Janse M J, Meijer F L and Arzbacher R C. Total excitation of the isolated human heart. *Circulation* 41:899 1970
- 4 Blondeau M and Lenegre J. Bloc atypique de la branche droite. Paris 1970. Masson et Cie
- 5 Lenegre J. Personal communication

Pork and hypertension

Several years ago it was indicated that pork will produce arterial hypertension.¹ This message has been ignored. Careful medical histories usually reveal that patients themselves learn that eating

pork will increase their blood pressure, produce headaches of varying severity, giddiness, scotomata, dyspepsia, pounding of the heart, beat, tinnitus and other symptoms and signs of rather sudden eleva-

the marked difference between ^{125}I fibrinogen-detected thrombi and the clinically detectable disease that some additional hypothetical factor is necessary for the former to develop into the latter. If this were so then it is highly probable that any treatment which prevented the formation of ^{125}I fibrinogen-detectable thrombi would also prevent the superimposed clinical disease. But one could also argue at least theoretically that it might be possible to block the hypothetical factor necessary for the development of the full clinical picture without necessarily altering the incidence of ^{125}I detectable thrombosis.

It is highly unlikely that aspirin would affect such a hypothetical additional factor and indeed the evidence is against it; nevertheless there are some who would argue that it would be desirable despite the discouraging findings in the present trial to carry out a further investigation involving very much larger numbers of patients to decide whether aspirin has an effect on clinical postoperative venous thrombosis or not, particularly since one study has been reported claiming to demonstrate some clinical benefit from the administration of aspirin.³

Clearly many factors both known and unknown contribute to the development of venous and arterial thrombosis. Local precipitating abnormalities must be important in thrombus formation because venous and arterial systems as a whole never clot solid. Platelets almost certainly play a far larger part in arterial thrombosis and it would seem rational indeed mandatory in view of the great importance of arterial thrombosis to evaluate the effect of aspirin in as many thrombotic situations as possible with a view to identifying those conditions where its specific effects have therapeutic benefit.

There is already convincing evidence that aspirin is of real value in patients with amaurosis fugax.^{4,5}

One recognizes daunting logistic problems in organizing suitable trials especially in myocardial infarction which is a rare disease in terms of patient years. Furthermore it will prove difficult to keep the control subjects from taking aspirin as it is now consumed on such a vast scale throughout the Western world. Nevertheless we hope such trials will be pushed through to their logical conclusion despite the evidence that aspirin is without effect in postoperative venous thrombosis and we await the results of those trials already under way with keenest interest.

J R O'Brien
St Mary's Hospital
Portsmouth PO3 6AG
W J H Butterfield
University of Nottingham
Nottingham NG7 2RD England

REFERENCES

- 1 Report of the Steering Committee of a trial sponsored by the Medical Research Council. Effect of aspirin on postoperative venous thrombosis. *Lancet* 2:441 1972.
- 2 O'Brien J R, Gulevski V and Etherington M. Two in vivo studies comparing high and low aspirin dosage. *Lancet* 1:1399 1971.
- 3 Salzman E W, Harris W H and De Sanctis R W. Reduction in venous thromboembolism by agents affecting platelet function. *N Engl J Med* 281:1287 1971.
- 4 Harrison M J, G Marshall J, Meadows J C and Ross Russell R W. Effect of aspirin in amaurosis fugax. *Lancet* 2:743 1971.
- 5 Evans G. Use of platelet suppressive drugs on the incidence of recurrent venous thrombosis and transient cerebral ischaemia. *Abstr Int Soc Thromb Haemostasis* Washington 40 1972.

Left hemiblocks revisited from the histopathological viewpoint

The introduction of the concept of left hemiblocks^{1,2} undoubtedly represented a fundamental step forward in the understanding of intraventricular conduction disturbances. The assumptions which formed the basis of this concept were extremely useful especially for didactic purposes. Nevertheless in course of time it seems that these assumptions deserve further attention in particular from the anatomical and histopathological viewpoints.

It is for example generally agreed that the left branch is anatomically a bifascicular system which acts physiologically as such.^{3,4} We recently had the opportunity to test this assumption in a histological study of the left bundle branch system carried out on 20 hearts from patients devoid of conduction defects. This investigation confirmed the consistent

presence of a thin and elongated anterior radiation and of a wider posterior one. However in addition to these two well known fasciculi the left branch was observed to frequently give off a third radiation designated to cover the midseptal surface (11 out of 20 cases). This easily identified structure emerged either from the common left bundle (five cases) or from the anterior (three cases) or posterior radiation (three cases). In the remaining nine instances the septal coverage was supplied either by the posterior radiation (three cases) or by a complicated plexus of ramifications given off by both the anterior and posterior fasciculi (six cases). The anatomical features were such that in most cases it seemed reasonable to describe the left ventricular Purkinje network as composed of three main widely inter-

Letters to the Editor

Digoxin tablets A possible problem with biological availability

To the Editor

The Food and Drug Administration suggests that pharmacists keep for the information of physicians a record of the brand name and manufacturer or supplier as well as the lot number of all digoxin tablets dispensed.

In a recent editorial in the *Journal of the American Medical Association*¹ FDA affirmed the possibility of variance in digoxin tablets and recommended particular vigilance when patients obtain a new supply whether made by the same or a different manufacturer.

FDA indicated that currently available information suggests the following:

1 Digoxin tablets from different manufacturers and suppliers may not be equally bioavailable; therefore may not be therapeutically equivalents.

2 If unexpected difficulty is encountered in digitalization with digoxin tablets the physician should consider bioavailability as a possible source of the problem.

The physician should be alerted to these two points when evaluating the status of his patients particularly at a time when they obtain a new drug supply whether of a second brand or even the same brand since the possibility of lot-to-lot bioavailability inconsistency cannot be ruled out at this time.

The FDA emphasized that if a bioavailability problem does exist with regard to tableted digoxin products it is because we have become more sophisticated in our ability to define the factors that influence drug therapy, rather than any identifiable change on the part of the industry.

The Agency pointed out that the systematic testing and voluntary certification programs instituted by the FDA in 1970 to insure that digoxin tablets contain the specified quantity of active ingredient and meet other compendial requirements do not address the problem of bioequivalence and that additional studies are underway to determine whether an actual problem exists and to determine its magnitude if it does exist.

For readers who desire detailed information about the problem of bioavailability of orally administered digoxin as presently understood, several articles are recommended.²⁻⁷

Additional pertinent information on the subject when it becomes available will be published in the *FDA Drug Bulletin*. Physicians and pharmacists who are not now receiving the *Bulletin* may do so by writing:

Assistant to the Director for Medical Communications
Bureau of Drugs (BD 40)
Food and Drug Administration
5600 Fishers Lane
Rockville, Md 20852

REFERENCES

- 1 Biologic availability of digoxin tablets. *JAMA* 224:243 1973.
- 2 Lindenbaum J, Mellow M H, Blackstone M O and Butler V P Jr. Variation in biologic availability of digoxin from four preparations. *N Engl J Med* 283:1344 1971.
- 3 Shaw T R D, Howard M R and Hamer J. Variation in biological availability of digoxin. *Lancet* 2:303 1972.
- 4 Huffman D H and Azarnoff D L. Absorption of orally given digoxin preparations. *JAMA* 222:957 1972.
- 5 Wagner J G, Christensen M, Sakmar E, Blair D, Yates J D, Willis P W, Sedman A J and Stoll R G. Equivalence lack in digoxin plasma levels. *JAMA* 224:199 1973.
- 6 Falch D, Teven A and Bjerkelund C J. Comparative study of the absorption, plasma levels and urinary excretion of the new and old Lanoxin. *Br Med J* 1:695 1973.
- 7 Lindenbaum J, Preibisz J J, Butler V P Jr and Saha J R. Variation in digoxin bioavailability: a continuing problem. *J Chronic Dis* (In press).

tions in arterial blood pressure. The substance in pork responsible for these reactions remains unknown. Some people are so sensitive to pork that mere traces of bacon grease or of pork seasoning in food are sufficient to increase arterial blood pressure to dangerous levels. Daily use of pork in food will maintain elevations in blood pressure in patients with arterial hypertension. Acute elevations in blood pressure will return to normal when the patients are placed on a pork free diet. Elimination of pork from the diet of most patients with so-called essential hypertension will render the regulation of blood pressure not only possible but easy.

The history of Mrs. V. S., a 42 year old Negro woman, illustrates the relation of pork intake to arterial hypertension. This patient develops a rather sudden elevation in arterial blood pressure when she eats pork chops (her favorite meat), develops a splitting and unbearable headache, blurring of vision, marked giddiness, pounding of her heart, tinnitus, and a feeling of being seriously ill and near collapse. Her blood pressure increases to 240/140 or more at rest on occasions after eating pork for a few successive days. These symptoms will last for hours and gradually subside if she abstains from pork. All clinical manifestations of illness disappear and her clinical state returns to normal including her blood pressure without medication.

This patient's 35 year old brother, B. M., participated in a family feast in which a hog was killed and fresh pork was consumed in large quantities. He

suddenly developed a malignant type of hypertension (220/170) with hematemesis, hematuria, blood in his stools, palpitation, hyperhidrosis, markedly impaired vision with the syndrome of hypertensive encephalopathy and coma, and then died quickly.

The magnitude of sensitivity to pork varies among people and the amount and days of consumption required to produce hypertension also vary among patients. Some patients manifest no reaction, whereas others have fulminating reactions as described for the two patients above. There is no doubt that a complete elimination of pork in all forms is a good dietary therapeutic rule in the management of arterial hypertension in all patients. In addition, the elimination of all forms of pork from the diet of all people may be a good practice for the prevention and the control of hypertension, one of the most common and important illnesses of man. Salted pork is even more hazardous to the health of man.

George E. Burch, M.D.
Department of Medicine
Tulane School of Medicine
New Orleans, La.

REFERENCE

1. Burch, G. E., Phillips, J. H. and Wood, W. The high pork diet of the Negro of the Southern United States (Editorial). *A. M. A. Arch. Intern. Med.* 100: 859, 1957.

Books received

✓ THE CLINICAL DELINEATION OF BIRTH DEFECTS
Edited by Daniel Bergsma M D Baltimore 1972
The Williams & Wilkins Company 325 pages
Price \$29 00

✓ CARDIOVASCULAR PHYSICAL DIAGNOSIS Edited by
Martin J Frank M D and Sergio C Alvarez Mena
M D Chicago 1973 Year Book Medical Publishers
Inc 186 pages

CLINICAL SCALAR ELECTROCARDIOGRAPHY Edited by

✓ Bernard S Lipman AB M D FACP FACC
Edward Massie AB M D FACP FACC
and Robert E Kleiger BA M D Chicago 1972
Year Book Medical Publishers Inc 721 pages

✓ AN ABC OF MODERN IMMUNOLOGY By E J Hol
borow M D Cantab M R C P M E C Path
Boston 1973 Little Brown & Company 94 pages
Price \$4 95

Book reviews

✓ **CHRONICALLY IMPLANTED CARDIOVASCULAR INSTRUMENTATION** Edited by Ernest P. McCutcheon New York and London 1973 Academic Press Inc. 482 pp. Price \$14.50

This book on the proceedings of a symposium held in Lexington, Kentucky, during October 17 to 20, 1971, on implanted cardiovascular instrumentation should interest physiologists and others studying cardiovascular physiologic phenomena. The many contributors and contributions review extensively the many problems of instrumentation techniques, data analysis, and application of implanted devices to study cardiovascular physiologic phenomena. Even though this book will not interest clinicians, the problems under study eventually will have clinical significance. Engineers, physicians, bioengineers, and biophysicists, as well as physiologists, will find this book to be extremely valuable. The reviews of techniques and application reveal the many complex and important cardiovascular problems yet

to be studied. This is a good and important publication.

✓ **CARDIAC ARRHYTHMIAS: The Twenty-Fifth Hahnemann Symposium** Edited by Leonard S. Dreifus, M.D., and William Likoff, M.D. New York 1973 Grune & Stratton Inc. 681 pp. Price \$29.75

With great interest in arrhythmias these days, this publication of the papers presented at the 25th Hahnemann Symposium should interest all cardiologists and internists. The many contributors review the arrhythmias for the reader. Of course, it contains nothing new, but anyone who wishes to study arrhythmias will find this to be a valuable publication. This publication brings up-to-date the discussions of the 14th Hahnemann Symposium on arrhythmias of 7 years previous. This book is recommended for the clinician. Like the previous publication on arrhythmias, this volume is presented from a practical point of view.

Acknowledgment to reviewers

The Editors wish to express their thanks and appreciation to the following who have aided in the review of manuscripts during the past year

| | | |
|-----------------|--------------------|------------------|
| F M Abboud | W Freyburger | J J Leonard |
| W H Abelmann | E D Frolich | H D Levine |
| R F Ackerman | S Furman | H J Levine |
| C M Agrest | Jack Geer | A J Liedtke |
| H E Aldridge | R W Gifford Jr | William Likoff |
| C S Alexander | Thomas Giles | J R Logic |
| J K Alexander | L Goldberg | R F Lowe |
| K Amplatz | Allan Goodyer | A A Luvada |
| G J Anderson | W J Grace | H A Lyons |
| B M Baker | H D Green | F G McMahon |
| S S Barold | D G Greene | D G McNamara |
| D V Bates | R L Grissom | O Magidson |
| A C Beall | R M Gunnar | D Mair |
| A A Berenbaum | J C Gunnells Jr | G V Mann |
| S Bernstein | A C Guyton | H J L Marriott |
| L F Bishop | V C Hall | Enrique Martinez |
| H Blackburn | R I Hamby | Daniel Mason |
| T M Blake | Jaak Han | D T Mason |
| D Blankenhorn | E W Hancock | Rashid A Massumi |
| R F Bond | J C Harkin | A M Master |
| R O Brandenburg | D C Harrison | H P Mauck Jr |
| E Breider | R M Harvey | J T Mazzara |
| A N Brest | W P Harvey | M Mendlowitz |
| I L Bunnell | R A Helm | A J Merrill |
| C A Caceres | M V Herman | J H Mitchell |
| E Carvajal | G R Herrmann | Yves Morin |
| Te Chuan Chou | I Hoffman | J J Morris Jr |
| J N P Davies | S W Hoobler | A G Morrow |
| R W DeSanctis | L G Horan | A S Nadas |
| K B Deser | M J Hughes | C V Nelson |
| H W Dhurandhar | H N Hultgren | C M Nice Jr |
| R Dillenkoffer | J O Neal Humphries | J A Noonan |
| H T Dodge | T N James | J Nyboer |
| Ephraim Donoso | L E January | R E Olson |
| Philip Dow | John Johnson | W H Parsons Jr |
| J T Doyle | Steve L Johnson | M L Pearce |
| H P Dustan | C E Jones | Lyle Peterson |
| E E Eddleman | C R Joyner | Ruth Pick |
| J E Edwards | W E Judson | H V Pipberger |
| W C Elliot | W B Kannel | W H Pritchard |
| L B Ellis | M H Kaplan | W L Proudfoot |
| M E F Engle | William Kirk | J R Pryor |
| E H Estes Jr | C E Kossmann | Elhot Rapaport |
| F A Finnerty Jr | J S LaDue | R J Reed |
| James Fisher | L E Lamb | E W Reynolds Jr |
| Nancy C Flowers | Richard Langendorf | J L Reynolds |
| N O Fowler | P H Langer | L W Reynolds |
| E D Fres | Louis Lemberg | Kay Rives |

Announcements

Cardiology symposium

The University of Texas Health Science Center at Houston Division of Continuing Education will present a Cardiology Symposium on December 3 through 6 1973 at Houston Texas This program will present an intensive review in cardiology The guest lecturer will be James J Leonard M D Professor and Chairman Department of Medicine School of Medicine University of Pittsburgh Pittsburgh Pa

For further information write The Office of the Director The University of Texas Health Science Center at Houston Division of Continuing Education P O Box 20367 Houston Texas 77025

Eleventh Annual Cardiology Seminar

The Rogers Heart Foundation announces the eleventh annual Cardiology Seminar under its sponsorship to be held at the Princess Hotel Southampton Bermuda on December 6 through 9 1973 The seminar theme is Coronaries and Controversies Program director is Henry J L Marriott M D Faculty are Dr Agustín Castellanos Miami Dr Robert Eliot Omaha Dr Desmond Julian Edinburgh Dr Frank LaCamera Jr St Petersburg Dr Harold Mankin Rochester Dr Henry J L Marriott St Petersburg Dr Victor Parsonnet Newark Dr Joseph Reeves Birmingham and Dr Leon Resnekov Chicago

For further details please write Rogers Heart Foundation St Anthony's Hospital St Petersburg Fla 33705 Telephone (813) 894 0790

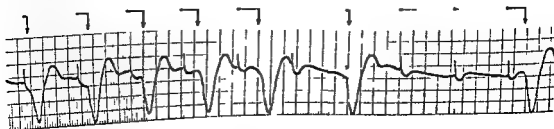
Annual meeting of American Thoracic Society

The Annual Meeting Committee American Thoracic Society Medical Section of the American Lung Association invites submission of papers on all scientific aspects of respiratory disease for presentation at the 1974 annual meeting in Cincinnati Ohio May 12 through 15 1974 Membership in the Society is not a prerequisite to participation in the program Abstracts of papers should be submitted to the Chairman of the Annual Meeting Committee before January 15 1974 Abstracts of papers from investigators outside the United States will be welcome provided they are in English and are submitted with the understanding that all papers accepted must be presented clearly in English at the Annual Meeting

Required forms and instructions may be obtained by writing Joseph F Tomashefski M D Chairman Annual Meeting Committee American Thoracic Society 1740 Broadway New York N Y 10019

Erratum

In the August 1973 issue of AMERICAN HEART JOURNAL in the article Mechanisms of cardiac arrhythmias From hypothesis to physiologic fact by Dr Alfred Pick Fig 35 on p 266 was inadvertently reversed The correct figure is illustrated below



An assessment of human cardiac transplantation

A Monim A Fadali MD*
Louis A Soloff MD**
Philadelphia Pa

On August 24 1972 Louis B Russel Jr an Indianapolis school teacher celebrated the fourth anniversary of his heart transplant. No one else has lived so long after heart transplantation.¹ His four year-old heart used to belong to a seven year-old boy from Providence Forge Virginia who was killed while stepping out of a restaurant during a ghetto fight.² Lower and his team performed the operation at the Medical College of Virginia. This outstanding achievement and the recent reports from Shumway's group^{3,4,5} at Stanford University make it important for those of us not actively participating in cardiac transplantation to take stock of what has been accomplished by and what has been learned from this operation. This knowledge is important to assess the place of cardiac transplantation in the overall management of the grossly disabled cardiac patient.

Transplantation of organs and tissues intrigued the imagination of mankind since antiquity. The blind⁶ thus described a chimera

A mingled monster of no mortal kind
Behind a dragon's fiery tail was spread
A goat's rough body bore a lion's head
Her pitchy no-trills flaky flames expire
Her gaping throat emits infernal fire

This mythical monster after devastating Caria and Lycia was killed by Bellerophon mounted on his winged steed Pegasus. The word chimera has recently been introduced into medicine to describe an animal which possesses for a long time viable cells from another animal.

The chimera was for centuries a figment of the imagination and the adjective chimerical became synonymous with imaginary. Nonetheless the universal significance of this concept is attested by its persistent influence on the arts of many civilizations from antiquity to the present day. Except for minor procedures on the face nose and penis the ancients had to be satisfied with prostheses. The Talmud⁷ records the use of crutches artificial limbs and teeth procedures that were practiced by other civilizations before and at that time. The increased need for prosthesis during the two world wars combined with the constantly developing industrial revolution led to the introduction of many sophisticated and electrically controlled prostheses which could simulate the movements of almost all voluntary and many involuntary groups of muscles including the fine movements of the thumb and fingers. All of the prostheses were applied to the

*In the Section of Thoracic and Cardiovascular Surgery and the Division of Cardiology Temple University Health Science Center Philadelphia Pa.

Support in part by The Council for Tobacco Research U.S.A. Inc.

Received for publication Oct 3 1972

Reprint requests to Louis A. Soloff MD Division of Cardiology Temple University Health Science Center 3401 N. Broad St. Philadelphia, Pa. 19140

Address reprint requests to Section of Thoracic and Cardiovascular Surgery Temple University Health Science Center
**Professor of Medicine Division of Cardiology Temple University Health Science Center

W Roberts
G G Rowe
H I Russek
J R Ryan
D A Rytand
Philip Samet
D C Scarpelli
J R Schenken
David Scherf
Leonard Scherlis
R M Schlant
Sidney Schnur
M L Segal
Ralph Shabetas
J T Shepherd
L Shewey

Maurice Sokolow
J F Spann Jr
Albert Starr
W J Stuckey Jr
Borys Surawicz
H J C Swan
W Jape Taylor
John Thomas
Bernard Towers
W G Walker
Richard Wasserburger
Yoshio Watanabe
W H Wehrmacher
M H Weil
S L Weinberg
L G Welt

S Wessler
W J Whalen
J P Whisnant
P D White
E D Wigle
Douglas Wilkerson
T Winsor
A C Witham
Norman Woody
R G Yaeger
Paul N Yu
R F Zelis
Harry F Zinsser
D P Zipes
M Ziskind
P M Zoll

An assessment of human cardiac transplantation

A Moneim A Fadal MD*
Louis A Soloff MD**
Philadelphia Pa

On August 24 1972 Louis B Russel Jr an Indianapolis school teacher celebrated the fourth anniversary of his heart transplant. No one else has lived so long after heart transplantation.¹ His four year old heart used to belong to a seven teen year old boy from Providence Forge Virginia who was killed while stepping out of a restaurant during a ghetto fight.² Lower and his team performed the operation at the Medical College of Virginia. This outstanding achievement and the recent reports from Shumway's group³⁻⁵ at Stanford University make it important for those of us not actively participating in cardiac transplantation to take stock of what has been accomplished by and what has been learned from this operation. This knowledge is important to assess the place of cardiac transplantation in the overall management of the grossly disabled cardiac patient.

Transplantation of organs and tissues intrigued the imagination of mankind since antiquity. The Iliad⁶ thus described a chimera

A mungled monster of no mortal kind
Behind a dragon's fiery tail was spread
A goat's rough body bore a lion's head
Her putchy nostrils flaky flames expire
Her gaping throat emits infernal fire

This mythical monster after devastating Caria and Lycia was killed by Bellerophon mounted on his winged steed Pegasus. The word chimera has recently been introduced into medicine to describe an animal which possesses for a long time viable cells from another animal.

The chimera was for centuries a figment of the imagination and the adjective chimerical became synonymous with imaginary. Nonetheless the universal significance of this concept is attested by its persistent influence on the arts of many civilizations from antiquity to the present day. Except for minor procedures on the face nose and penis the ancients had to be satisfied with prostheses. The Talmud⁷ records the use of crutches artificial limbs and teeth procedures that were practiced by other civilizations before and at that time. The increased need for prosthesis during the two world wars combined with the constantly developing industrial revolution led to the introduction of many sophisticated and electrically controlled prostheses which could simulate the movements of almost all voluntary and many involuntary groups of muscles including the fine movements of the thumb and fingers. All of the prostheses were applied to the

* In the Section of Thoracic and Cardiovascular Surgery and the Division of Cardiology Temple University Health Sciences Center Philadelphia, Pa.
Supported in part by The Council of Tobacco Research USA Inc.
Received by JAMA Oct 3 1972.
Reprinted from: Annals of the New York Academy of Sciences, Vol. 257, No. 1, pp. 1-10, 1980.
Address reprint requests to Dr. Soloff, Division of Cardiology, Temple University Health Sciences Center, 3401 N. 32nd St., Philadelphia, Pa. 19140.
Address correspondence to Dr. Fadal, Section of Thoracic and Cardiovascular Surgery, Temple University Health Sciences Center, 3401 N. 32nd St., Philadelphia, Pa. 19140.

external surface of the body. Except for a few botanical¹ and dental procedures, rare bone, thyroid, and experimental kidney transplant^{2,3} internal prostheses and chemical operations had to await the development of a variety of apparently unrelated techniques and the recognition of their applicability to medicine. Some of these techniques are instruments for cutting, anatomic dissection, experimental physiology, anesthesia, intubation, roentgenology, blood and tissue typing, heparin and cellophane, all eloquent testimony to the anticipated dividends of basic and industrial research to the welfare of mankind.

We cannot resist speculating on the future of the artificial heart because its future will we think greatly influence the usefulness and perhaps the practice of cardiac transplantation. Of all the internal organs, the heart, arteries and veins appear the most likely candidates for prosthetic replacement. This is so because the vital function of the heart is to act as a pump and that of the arteries and veins to convey blood. Pipes and pumps are well known products of industry. There is no difficulty in making and energizing a pump powerful enough to propel the amount of blood adequate to meet the needs of the body and in making a pipe strong enough to withstand the flow of blood. That the blood could tolerate so well a large clumsy foreign body like the Starr Edwards valve was in astonishing revelation. This success helps to sustain our faith in the ultimate development of a successful artificial heart. At the present time, Kolff⁴ states that the limiting factors for total cardiac replacement are thrombosis, hemolysis, and right heart failure. Dogs with an artificial heart have lived as long as eight days. From the standpoint of survival, these dogs are equivalent to Mann and colleagues⁵ dogs with transplanted hearts in 1933. We have every reason to believe that continued collaboration between biological and physical scientists with adequate facilities and financial support can solve this important problem in less than the 35 years that elapsed between Mann's experiment and the first successful human cardiac transplantation. Such an achievement would be of immense importance. It would solve the problems of

supply and immunologic complications, two main obstacles hindering practicability of cardiac transplantation and would also obviate some difficult and distasteful ethical questions. At the present time, from the standpoint of survival, the artificial heart is not a reality. Cardiac transplantation is

Correl and Guthrie¹⁰ in 1905 transplanted a heart to the neck of a dog. The heart beat spontaneously for one hour. Using the same model, Mann and associates⁵ in 1933 reported survival periods up to eight days with normal heart beats. Faced with a patient in terminal heart failure and with no suitable human donor available, James D. Hardy¹¹ of Jackson, Mississippi, transplanted the heart of a large chimpanzee to the dying patient on January 23, 1964. The transplanted heterograft maintained a blood pressure of 50 to 90 mm Hg for approximately one hour. The first human heart homotransplant was performed by Christian Barnard¹² of South Africa on December 3, 1967. This accomplishment aroused intense controversy, varying from absolute praise expressed by J. E. Edwards¹³, President of the American Heart Association, who called the procedure "a courageous and new advance in cardiology" and a milestone in the history of medicine and surgery, to utter condemnation and anger expressed by the Nobelist Wilhelm Lörsmann¹⁴ who compared heart transplantation to some of the Nazi experiments on humans. Enthusiasm was contagious and in the following year a total of 101 heart transplants were performed in several countries. Enthusiasm waned the following year because of the unacceptable mortality rate and possibly also for ethical and legal reasons. As of August 1, 1972, 194 heart transplants have been performed in 21 countries by 61 teams (Table 1). At least two of these patients had combined heart and lung transplantation because of associated severe pulmonary hypertension in one and advanced chronic obstructive pulmonary disease in the other.¹⁵ Two other patients underwent a second transplantation due to chronic rejection and graft failure after ten and seven months but both died in the early postoperative period.¹⁶ Twenty-seven recipients are current survivors from two to 45 months after

Table I Human heart transplantation—Chronology and world distribution (Dec 3 1967 to Aug 1 1972)*

| Year | World totals | U S A | Canada | France | South Africa | Other countries |
|-------|--------------|----------|--------|--------|--------------|-----------------|
| 1967 | 2 | 1 | 0 | 0 | 1 | 0 |
| 1968 | 101 (4)† | 54 (4) | 14 (2) | 10 (1) | 2 | 21 |
| 1969 | 47 (3) | 34 (2) | 1 | 0 | 4 (1) | 8 |
| 1970 | 17 (5) | 16 (4) | 1 (1) | 0 | 0 | 0 |
| 1971 | 17 (7) | 13 (6) | 1 | 0 | 3 (1) | 0 |
| 1972‡ | 10 (5) | 8 (5) | 0 | 0 | 1 | 1 |
| Total | 194 (27) | 126 (21) | 17 (3) | 10 (1) | 11 (7) | 30 |

American College of Surgeons and the International Health Organization Registry
†Number in parentheses indicate number of cases.
‡Data not yet complete.

Table II Human heart transplantation—Prolonged survival*

| | > 6 mos | > 1 yr | > 2 yr | > 3 yr | > 3½ yr |
|---------------------------------------|---------|--------|--------|--------|---------|
| Total recipients 191 | 49 | 36 | 21 | 11 | 9 |
| Current survivors 27 as of Aug 1 1972 | 24 | 19 | 14 | 9 | 7 |

American College of Surgeons and the International Health Organization Registry

transplantation nine of them living over three and one half years (Table II). The majority of the transplants were performed with ischemic cardiac arrest of the donor's heart according to the technique standardized by Lower and associates.¹⁰ Operative deaths were extremely rare and most of the mortalities occurred in the first three months after surgery. The attrition rate declined sharply after the third postoperative year. The overall six month one year two year three year and three and one half year survival rates are presently 70 per cent 17 per cent 11 per cent 7 per cent and 47 per cent respectively. The group at Stanford University California headed by Dr Norman Shumway who have dedicated themselves to the study of this problem obtain much better results. The overall survival rate in their series was 37 per cent at one year and 26 per cent at two years.¹¹ The results of this group show steady improvement. In 1968 22 per cent of their transplant patients survived one year whereas 30 per cent of those operated upon in 1970 to May 1971 survived

at least one year.¹² Almost all the long term survivors had achieved impressive degrees of physical social and vocational rehabilitation. It is therefore clear that patients can survive cardiac transplantation and improve sufficiently to lead a useful life. Indeed in the hands of the experienced cardiac surgeon the operative procedure no longer significantly limits survival. However the percentage of patients who improve sufficiently to lead a useful life for an appreciable length of time is still disappointingly small.

At first a great deal of effort was spent on determining whether the cardiac transplant could be reinnervated and if not on the denervated heart capacity to perform work. This is surprising in view of a classic experiment performed by Patterson Piper and Starling¹³ in 1914. They showed that the denervated isolated heart possesses an intrinsic mechanism by which it can meet the demands of increased pressure and volume. Casser and Meek¹⁴ in the same year and Saman¹⁵ in 1935 did produce evidence that the dog's ability to exercise

was impaired by denervation. Their experiments were criticized by other observers²⁰ who showed by employing similar techniques that other organs were also denervated and this may have altered the results. Finally Donald and Shepherd^{1,2} demonstrated that the Frank-Starling mechanism was operative in dogs with chronic cardiac denervation. Blinks³ demonstrated that in the isolated heart distension of the right atrium had a positive chronotropic effect in several animal species. However, Donald and Shepherd⁴ could not find a linear correlation between increments of right atrial pressure and rises in heart rate in dogs with chronic denervation. The most plausible explanation for the increase in heart rate and at least part of the increase in cardiac output is the action of circulating catecholamines.⁶ Denervation depletes the myocardial stores of catecholamines²⁷ and this renders the heart more sensitive to the action of circulating catecholamines. This can explain the observed faster heart rate of the allograft heart at rest in comparison to control subjects. The catecholamines released during exercise will also be effective. In one patient²⁸ several months after heart transplantation measurement of the blood level of epinephrine and norepinephrine during exercise showed significant rise synchronous with the observed increase in heart rate. Actually, immediately after operation the heart resumes function either spontaneously or after electrical stimulation.

There is overall agreement on the hemodynamic observations after human cardiac transplantation. The reports from Shumway's group⁹ are the most impressive because their findings were obtained one and two years after operation and while their patients were clinically stable.

In essence at rest the heart rate is slightly faster than normal and not responsive to respiration, the intracardiac pressure is normal, the cardiac index is low normal or just below normal, the stroke index is slightly below normal and the arteriovenous oxygen difference is slightly greater than normal. This last form of failure suggests that the patient would not be able to tolerate exercise for long, if at all, which is maximal for the normal. Indeed, this group

found that arterial blood analysis during submaximal exercise showed evidence of anaerobic metabolism at relatively low work loads and levels of oxygen uptake. This occurs even though the ratio between an increase in cardiac output and an increase in oxygen uptake during exercise remains normal.

During exercise the left ventricular end diastolic pressure rises abruptly, the heart rate increases slowly to a plateau and after exercise it takes 20 minutes or more to reach its resting level. The cardiac output increases without a significant change in left ventricular ejection time and the arteriovenous oxygen difference increases inordinately. The systemic and left ventricular pressures rise progressively for the first few minutes of exercise and then reach a plateau for the remaining five minutes of exercise.

It is noteworthy that the pulmonary artery pressure and resistance after transplantation can be close to normal. That this is not always true is indicated by two of the first ten deaths reported by this group.

The precise causes of the hemodynamic abnormalities are uncertain. Unfortunately the chronologic clinical history of each patient is not given in most reports. Although at least some of these findings resemble those in the canine denervated heart and in the chemically denervated human heart, one cannot exclude the effects of increased afterload, chemotherapy used at the time of study, and the effects of rejection most likely present at least in the past. The last effects are suggested by the marked deterioration at two years of one patient who at that time revealed severe diffuse narrowing of all the major coronary arteries with complete obliteration of the proximal left circumflex artery. Such arterial changes can occur without clinical evidence of rejection. It is this rejection phenomenon and the untoward effects of treatment aimed at preventing and reversing it that pose the stumbling block that must be removed before cardiac transplantation can pass from an experimental procedure to a clinically acceptable method of treatment.

The rejection reaction is not related to the recipient's type of heart disease. Similar lengths of survival have been reported

after operations on patients with cardio myopathy congenital rheumatic and coronary heart disease and even in one patient with rheumatic heart disease and pulmonary artery pressure at systemic levels. Transplantation therefore appears to be equally suitable for all types of heart disease and would be contraindicated only in those patients with extracardiac disorders that seriously limit life and also by absolute biologic incompatibility. Finally none of the above dysfunctions of the transplanted heart have been of a sufficient magnitude to preclude a useful life and therefore would not be of any significant concern if further deterioration did not occur.

Unfortunately present day methods of diagnosis and treatment have not been able to prevent the development of the rejection reaction. Absolute contraindications to the use of a donor's heart for a particular recipient are (1) ABO blood group incompatibility (2) preformed antibodies in the recipient's serum to donor's lymphocytes and (3) gross disparity in the size of the donor's and recipient's hearts. Rejection has occurred in the absence of other demonstrable incompatibilities and survival has been reported in instances with gross mismatches.¹¹ These results suggest the likelihood of unrecognized factors such as chemically impure antigens and specific tissue loci not manifested by present day serologic tests. Dr Amos of Duke University and Dr Bach of the University of Wisconsin¹¹ have recognized such leucocyte-defined loci and hopefully a practical test for their clinical recognition will be forthcoming in the near future. In the meantime common sense dictates a choice of a donor with minimal antigenic disparities for a specific recipient.

As long as the rejection reaction cannot be prevented the best way to cope with the situation is to detect it early and to treat it properly with agents that have the least untoward effects. Such agents are also used with the hope of preventing postponing and attenuating the rejection reaction.

Early detection is suggested by electrocardiographic (ECG) evidence of decreased voltage, atrial arrhythmias and rightward deviation of the QRS complexes.¹² In some patients the appearance of a diastolic

gallop was the first sign of rejection preceding the ECG changes.¹³ DeBakey and colleagues¹² reported that symptoms of anorexia and malaise can also precede ECG changes by 1 to 16 hours.

Roentgenologic and ultrasonic evidence of increasing thickness of the left ventricular wall, increased size of the right ventricle and the whole heart and changes in cardiac contractility have all been helpful.¹² Clinical signs of cardiac insufficiency and the rise in enzymes (CPK, LDH) are usually late signs of rejection. More recently the Stanford group¹⁴ has been doing endocardial biopsies on their patients to obtain baseline data with respect to the expected changes in the event of smoldering and chronic rejection.

Evidence is accumulating that immunologic abnormalities precede and are more specific than all other evidences of rejection and can be used as a guide to intensify treatment with the hope of preventing clinical disability. One method consists in serial measurements of serum titers of fluorescein tagged anti IgG globulins. They are bound primarily to the sarcolemmal sheath of the myofibers.¹⁵ Another method is serial quantification of the level of inhibition of the conversion of C₃.¹⁶ A third method is serial determination of the rate of spontaneous transformation of blood lymphocytes to blast cells in culture.¹⁷ These methods appear more sensitive than myocardial scanning with ¹³¹cesium.¹⁸

Azathioprine (Imuran) and prednisone are the drugs universally accepted for the prevention and treatment of rejection. The doses initially must be high yet carefully regulated because both drugs enhance susceptibility of the host to pathogenic and opportunistic organisms. The effectiveness of other methods of treatment has not as yet been established.

Most recipients have received anti lymphocytic globulin (ALG) or a purified globulin preparation from horse antihuman thymocyte serum for several weeks or months after operation. Controversy concerning their value may be related to the development of antibodies against the foreign protein and the fact that these globulins are eliminated from the circulation 8 to 12 days after treatment.¹⁹ Nonetheless a

short term course may be of value in an emergency, particularly when leukopenia and hepatic dysfunction might contraindicate the use of Actinomycin D and increased doses of Imuran. There is evidence that ALG might enhance tumor growth.³⁹ Mice given ALG could not develop immunity to a second implant of sarcoma 180. However, none of the animals developed a spontaneous tumor.⁴⁰ At the Cleveland Clinic⁴¹ 65 out of 200 recipients of renal transplants were given intramuscular injections of horse anti human ALG. In one case a reticulum cell sarcoma developed at the site of the intramuscular injection of ALG. However, the possible ALG potential for induction of neoplasia does not at this stage justify discontinuing this agent in the treatment of rejection.

Another method of treatment is based on findings in rats after skin allografting.⁴² During periods of rejection it was found that the urinary histamine excretion was increased and also the histamine forming enzyme histidine decarboxylase activity in the allograft increased by 33 fold during rejection. Administration of histidine decarboxylase inhibitors prolonged the survival of skin allografts between inbred rats and suppressed the formation of antibodies to antigens.⁴³ The best results were achieved when semi carbazide and diamine hydrazide was combined with a pyridoxine deficient diet.⁴⁴ A close chemical relation exists between semi carbazide and isonicotinic acid hydrazide (INH). This group at UCLA Harbor General Hospital has added INH and histadyl H an antihistamine to the standard regimen of Imuran and prednisone. The program was started at operation in four renal and one heart transplant patients. None of them showed evidence of acute rejection up to six months after operation despite the fact that all were Terasaki C or D matches.

Local graft irradiation which has been found valuable after experimental and clinical renal homotransplantation is an attractive idea because it circumvents the systemic side effects of immunosuppressants. Irradiation injury to the myocardium is probably unlikely because the suggested doses are comparatively low. However, an underlying immunologic injury might en-

hance irradiation injury. The value of local graft irradiation after canine cardiac allografting has been shown by Graham and associates.⁴⁵ In 60 per cent of the animals serial myocardial biopsies showed striking histologic improvement when local radiation was employed to modify acute rejection episodes. The mechanisms involved are not entirely clear. The efferent arc of rejection might be interfered with through destruction of the lymphocytes which are known to be radiosensitive.⁴⁶ There is also evidence that the efferent arc or sensitization of the host might be impaired by graft irradiation.⁴⁷ Improved graft perfusion has been noted after irradiation due to amelioration of tissue edema associated with rejection.

The role of the thymus has been fascinating to those involved in organ transplantation. The cell mediated immune response rests on the long-lived small lymphocytes. They are known to be thymus dependent. After a mysterious voyage from the bone marrow to the thymus gland, the small lymphocytes become immunologically competent.⁴⁸ Despite this crucial role of the thymus its removal at the time of transplantation apparently did not improve graft survival in clinical experience.

Perhaps the most successful approach to the problem of organ rejection will be the induction of immunologic enhancement.⁴⁹ Immunologic enhancement means the prolongation of allograft survival by the presence in the recipient of an alloantibody directed against the alloantigens of the donor's graft. The process may be accomplished by active immunization of the graft recipient with donor tissues or passively by administering to the recipient serum containing alloantibodies. This alloantisera is prepared by immunizing another individual. It is not xeno or heteroanti serum as is the case with ALG. The alloantibodies by destroying or blocking the graft antigens will eliminate the destructive cellular responses to the graft antigens and therefore graft survival is prolonged. The greatest advantage of immunologic enhancement is its specificity. It does not increase the recipient's susceptibility to infection or malignancy as noted with modalities that are used to induce immunologic tolerance such as immunosuppression and radiation.

The best results were achieved when active and passive immunization were combined.

According to Nossal⁴² if the rules governing the activation of killer cells versus antibody forming cells could be described one can guarantee tilting the balance both in cancer and in transplantation.

All these measures are most effective in suppressing the early rejection phenomenon. This early rejection is characterized anatomically by edema, endothelial swelling, sloughing and inflammatory changes. However, these changes are not always manifested clinically and by present laboratory tests. Furthermore, clinical reversibility does not exclude permanent anatomic damage. Shumway^{30,32} agrees that the separation of reversible from irreversible morphology during rejection is not clear. This chronic form of rejection is not responsive to the measures listed above. It is characterized anatomically by intimal proliferation with obliterative coronary artery disease and can begin as early as nine days after operation and be completely silent until gross and irreversible myocardial dysfunction occurs. Thus in three patients whose grafts showed the most severe signs of rejection, numerous ECGs taken at rest and during exercise, ballistocardiograms, ultrasonic cardiograms and other tests failed to be diagnostic of rejection. Therefore, it may be assumed that the usual immunosuppressive drugs were not effective in preventing this serious complication.

More recently, Shumway and colleagues⁴³ have reported normal coronary arteries as visualized by coronary arteriography in three patients studied one year after operation. This group attributes the good success in the 3 patients to the use of a low cholesterol, low saturated fatty acid diet and to the use of warfarin and dipyridamole originally suggested by Kahn and associates⁴⁴ and by Kincaid Smith.⁴⁵ These findings are extremely encouraging but their significance is not clear because there was no change in the blood cholesterol. Coronary arteriography does not exclude gross atherosclerosis and the time interval is too short. This group had already shown that one cannot predict what changes will occur at two years based upon the findings at one

year after operation. Finally, the deleterious effect of residual pulmonary vascular disease on the function of the right ventricle may explain predominant right heart failure after cardiac transplantation.

Kosch and associates³¹ state that graft atherosclerosis resembles spontaneous atherosclerosis and any differences between the two can be explained by the differences in rates of development of the atherosclerosis. This would suggest that spontaneous atherosclerosis is an autoimmune disease. Such a possibility is supported partly by Beaumont's³³ finding that hyperlipoproteinemia may on occasion be due to an autoimmune response. It should be remembered however that the number of ways that a tissue can respond to injury is limited whereas the number of injurious agents is limitless. We have not had the opportunity of examining graft atherosclerosis. Acute rejection is described as involving capillaries, vasorum, veins and large vessels of the coronary arterial system. The chronic form is characterized by major involvement of the first branches of the intramyocardial portion of the coronary arteries and a tremendous proliferation of the intima. These findings do not appear to us to resemble spontaneous atherosclerosis.

Even after the rejection reaction will have been conquered, other major problems relating to cardiac transplantation remain. One is logistics relating to procurement and storage of cardiac allografts. Organ banks are badly needed for prolonged storage. Short methods of preservation are also required to reduce the inevitable ischemic damage which occurs preterminally before removal of the heart from the donor and which continues until graft revascularization by the host. Various methods have been employed to preserve the heart in a viable state. These include hypothermia, hyperbaric oxygenation, freezing, metabolic inhibitors, interim perfusion and the use of intermediary hosts. Moderate hypothermia is known to enhance organ viability. Deep hypothermia frequently leads to interstitial edema which may interfere with organ function after transplantation. Kondo and colleagues³ resuscitated canine hearts after 24 hours storage in saline at 4°C. Lower³⁴ successfully homotrans-

short term course may be of value in an emergency, particularly when leucopenia and hepatic dysfunction might contraindicate the use of Actinomycin D and increased doses of Imuran. There is evidence that ALG might enhance tumor growth.³⁹ Mice given ALG could not develop immunity to a second implant of sarcoma 180. However, none of the animals developed a spontaneous tumor.⁴⁰ At the Cleveland Clinic⁴¹ 63 out of 200 recipients of renal transplants were given intramuscular injections of horse anti human ALC. In one case a reticulum cell sarcoma developed at the site of the intramuscular injection of ALC. However the possible ALG potential for induction of neoplasia does not at this stage justify discontinuing this agent in the treatment of rejection.

Another method of treatment is based on findings in rats after skin allografting.⁴² During periods of rejection it was found that the urinary histamine excretion was increased and also the histamine forming enzyme histidine decarboxylase activity in the allograft increased by 33 fold during rejection. Administration of histidine decarboxylase inhibitors prolonged the survival of skin allografts between inbred rats and suppressed the formation of antibodies to antigens.⁴³ The best results were achieved when semi carbazide and diamine hydrazide was combined with a pyridoxine deficient diet.⁴⁴ A close chemical relation exists between semi carbazide and isonicotinic acid hydrazide (INH). This group at UCLA Harbor General Hospital has added INH and histidyl H in antihistamine to the standard regimen of Imuran and prednisone. The program was started in operation in four renal and one heart transplant patients. None of them showed evidence of acute rejection up to six months after operation, despite the fact that all were Terraki C or D matches.

Local graft irradiation which has been found valuable after experimental and clinical renal homotransplantation is an attractive idea because it circumvents the systemic side effects of immunosuppressants. Irradiation injury to the myocardium is probably unlikely because the suggested doses are comparatively low. However an underlying immunologic injury might en-

hance irradiation injury. The value of local graft irradiation after canine cardiac allografting has been shown by Craham and associates.⁴⁵ In 60 per cent of the animal serial myocardial biopsies showed striking histologic improvement when local radiation was employed to modify acute rejection episodes. The mechanisms involved are not entirely clear. The efferent arc of rejection might be interfered with through destruction of the lymphocytes which are known to be radiosensitive.⁴⁶ There is also evidence that the efferent arc or sensitization of the host might be impaired by graft irradiation.⁴⁷ Improved graft perfusion has been noted after irradiation due to amelioration of tissue edema associated with rejection.

The role of the thymus has been fascinating to those involved in organ transplantation. The cell mediated immune response rests on the long lived small lymphocytes. They are known to be thymus dependent. After a mysterious voyage from the bone marrow to the thymus gland the small lymphocytes become immunologically competent.⁴⁸ Despite this crucial role of the thymus its removal at the time of transplantation apparently did not improve graft survival in clinical experience.

Perhaps the most successful approach to the problem of organ rejection will be the induction of immunologic enhancement.⁴⁹ Immunologic enhancement means the prolongation of allograft survival by the presence in the recipient of an alloantibody directed against the alloantigens of the donor's graft. The process may be accomplished by active immunization of the graft recipient with donor tissues or passively by administering to the recipient serum containing alloantibodies. This alloantiserum is prepared by immunizing another individual. It is not xeno or heteroanti serum as is the case with ALC. The alloantibodies by destroying or blocking the graft antigens will eliminate the destructive cellular responses to the graft antigens and therefore graft survival is prolonged. The greatest advantage of immunologic enhancement is its specificity. It does not increase the recipient's susceptibility to infection or malignancy as noted with modalities that are used to induce immunologic tolerance such as immunosuppression and radiation.

J. L. Me 86
V. 11 6

18 years of age or older may donate all or part of his body for medical purposes to take place after death. This can be spelled out in a uniform donor card. In its absence the next of kin may act. Legislation adopted in Kansas⁶⁶ has spelled out the time of death as that time after all resuscitative measures have failed when there is either absence of spontaneous respiratory and cardiac function or an absence of spontaneous brain function. The ideal donor is one whose heart is still beating after spontaneous brain function has ceased. It is now agreed that the electroencephalogram should be flat for at least two hours in the normothermic patients and should not be one that is depressed by drugs. The act also provides that the time of death should be determined by physicians who attended the patient or who certified his death but these physicians cannot be those involved in removing and transplanting the part. We would add that death should be certified by a neurologist or neurosurgeon and by the electroencephalographer.

We hope that this definition of death will not discourage research on methods to revive an apparently dead brain. After all it was not so long ago when ventricular fibrillation and cardiac standstill were regarded as irreversible signs of death. Death of tissue is due to two independent and interacting factors: primary cessation of anabolism and secondary impairment and cessation of anabolism by a hostile environment. Obvious hostile environments are freezing and drugs. The catabolic effects of anabolism if not removed can by themselves choke and kill cells. These examples should encourage search for methods of determining the initial causes of brain damage and methods for reviving the brain.

Because of the frequency of occult coronary artery disease the donor should preferably be a young adult.

Who should be the recipient? It is a mistake to say that he should be the one who needs the heart most provided he has no other disorder that would limit the anticipated benefits from the transplant and provided he and preferably his family understand and accept the risks involved. The question however is not so simple because prognosis particularly for the

immediate future of those with coronary heart disease is not easy to define and because the demand far outstrips supply. The Shumway group¹⁸ is to be congratulated on the excellence of their diagnosis and prognosis. The average survival rate of 20 patients whom they selected for transplantation but could not be operated on for various reasons was 30 days. Half of them died by the tenth day after selection and only one patient lived more than 90 days. This survival rate is significantly lower than that of transplant recipients at the same institution.

It should be mandatory to preserve a chronologic detailed history, findings and treatment of all patients who are being considered for transplantation. Preference should be given to the younger patient but otherwise the physician should not try to judge the value of the life he hopes to save.

Considering all these unsolved problems is the operation worthwhile? We agree with Claude Bernard⁶⁷ who a century ago defined the limits of human experimentation.

It is our duty and our right to perform an experiment on man whenever it can save a life, cure him or gain him some potential benefit. The principle of medical and surgical morality therefore consists in never performing on man an experiment which might be harmful to him to any extent even though the results might be highly advantageous to science that is to say to the health of others. But performing experiments and operations exclusively from the point of view of the patient's own advantage does not prevent their turning out profitably to science.⁶⁸

Who then should perform the operation? We agree with Nossal¹⁷ that the research with respect to cardiac transplantation should proceed cautiously, remaining the province of the small number of groups with a profound commitment. These groups should be composed of scientists whose major research interest is in transplantation. They should have adequate institutional facilities and paramedical help. All persons engaged in this endeavor should be adequately supported so that if necessary they can devote their full time to the care of the patients. There should be no charge whatsoever either personal or institutional.

planted a dog heart after a hypothermic anoxic period of seven hours. Using hyperbaric oxygen at three atmospheres in conjunction with hypothermia, the dog's heart was preserved up to 72 hours in a viable condition. However Lyons and associates³⁶ achieved the same result with hyperbaric helium or nitrogen at three atmospheres, suggesting that pressure probably acts by inhibiting metabolism and preventing tissue edema.

Various methods and solutions have been used for perfusion with and without hypothermia in order to maintain organ viability. Methods utilized varied from the simple Langendorff columns³⁷ and the pulsatile perfusion system of Carrel and Lindbergh³⁸ to the elaborate heart lung preservation chamber designed by DeBakey and associates.³⁹ Crystallloid and colloid solutions, plasma and whole blood were used and a long list of additives was frequently recommended: insulin, hydrocortisone, magnesium sulfate, penicillin and other antibiotics, tris buffer and heparin—to mention a few. High osmolality of the perfusate and hyperbaria were found beneficial to prevent edema formation. However in any artificial system the perfused organ after a variable number of hours showed increased vascular resistance manifested by a rise in perfusion pressure, falling venous output and increasing edema.⁴⁰ Metabolic and functional deterioration subsequently became apparent. The duration of organ viability with artificial perfusion has not been firmly determined. With the present status of perfusion we think that 48 hours of organ preservation is a maximum beyond which successful organ transplantation would be highly unlikely. This period of time would allow function if evaluation of the preserved organ, tissue typing, and transportation of the heart to a recipient and perhaps immunologic enhancement by perfusion with specific antibodies.

Organ preservation by freezing has been unsuccessful so far. During standard slow freezing that is about 1° C per minute extracellular ice crystal formation occurs. Water is withdrawn from the cells to the extracellular ice crystals. Intracellular osmolarity rises and cellular injury results. To solve this problem cryo protective

agents and super cooling have been proposed. Cryo protective agents⁴¹ such as dimethylsulfoxide and glycerine tend to inhibit extracellular ice formation and therefore to decrease cellular dehydration. However these agents penetrate into the cells at a much slower rate than water and therefore during thawing osmotic damage to the cells might occur. With fast or super cooling intracellular ice crystal formation occurs and cell survival declines. Apart from these problems the main obstacle to freezing whole organs is that each population of cells has its own specific velocity of cooling. An optimum velocity for one kind of cell might be harmful to another kind.

Metabolic inhibitors were shown to prolong organ viability to at least double that of control organs.⁴² The most promising metabolic inhibitor is 2 per cent magnesium sulfate. In addition to slowing the metabolic rate it maintains membrane integrity and transmembrane potentials.⁴³ Intermediary transplantation in a temporary host has been suggested as a method of organ preservation by Angell and Shumway.⁴⁴ These hearts were shown by Wells and associates⁴⁵ to maintain an adequate pump function after a period of seven weeks. However possible rejection from two hosts might undermine the usefulness of this method.

A number of moral and ethical issues related to organ transplantation must be resolved in the minds of those who are involved with the procedure.

In the present state of the art of organ preservation the best chance of successful cardiac transplantation is the use of a donor heart that is still beating or has just stopped beating. However we should all heed these wise words of Ciertz: "A person dying is still a person living and he keeps his elementary human rights up to the moment when life is extinct."⁴⁶ There is a steady progression from clinical death to electrical (heart-brain) death to biologic death and then to cellular death which is temporally different in different parts of the body. When the brain is totally dead integrative vegetative life ceases with permanent extinction of body life.

The uniform anatomical gift act provides that any person of sound mind and

- tion by regional neural ablation. Description of the operation, verification of the denervation and its effects on myocardial catecholamines. *Circ Res* 9:275 1961
- 21 Donald D E and Shepherd J T. Response to exercise in dogs with cardiac denervation. *Am J Physiol* 202:393 1963
 - 22 Donald D E and Shepherd J T. Sustained capacity for exercise in dogs after complete cardiac denervation. *Am J Cardiol* 14:853 1964
 - 23 Blinits J R. Positive chronotropic effect of increasing right atrial pressure in the isolated mammalian heart. *Am J Physiol* 186:299 1956
 - 24 Donald D E and Shepherd J T. Changes in heart rate on intravenous infusion in dogs with chronic cardiac denervation. *Proc Soc Exp Biol Med* 113:315 1963
 - 25 Donald D E and Samueloff S L. Exercise tachycardia not due to blood borne agents in canine denervation. *Am J Physiol* 211:703 1966
 - 26 Angell W W, Dong E Jr and Shumway N E. A humoral substitute for nervous control in the dog heart transplant. *Surg Forum* 18:223 1967
 - 27 Cooper T, William V L, Jelinek N and Hanlon C R. Heart transplantation. Effect on myocardial catecholamine and histamine. *Science* 138:40 1962
 - 28 Leachman R D, Leatherman L L, Rochelle D G, Cooley D A, Hallman G L, Bloodwell R D and Nora J J. Physiologic behavior of the transplanted heart in six human recipients. Abstract presented at the Eighteenth Annual Scientific Session of the American College of Cardiology, New York, February 26 to March 2 1969
 - 29 Stinson E B, Schroeder J S, Griep R B, Dong E Jr and Shumway N E. Hemodynamic observations one and two years after cardiac transplantation in man. *Circulation* 35:1183 1972
 - 30 Griep R B, Stinson E B, Dong E Jr, Clark M A and Shumway N E. Acute rejection of the allografted human heart. Diagnosis and treatment. *Ann Thorac Surg* 1:113 1971
 - 31 Amos D H and Bach F H. Phenotypic expressions of the major histocompatibility locus in man (HLA). Leukocyte antigens and mixed leukocyte culture reactivity. *J Exp Med* 128:623 1968
 - 32 Stinson E B, Dong E Jr, Bieber C P, Jopp R L and Shumway N E. Cardiac transplantation in man II. Immunosuppressive therapy. *J Thorac Cardiovasc Surg* 58:3 6 1969
 - 33 DeBakey M E, Diethrich E H, Glick G, Noon G P, Butler W T, Rossen R D, Liddcoat J E and Brooks D K. Human cardiac transplantation. Clinical experience. *J Thorac Cardiovasc Surg* 58:303 1969
 - 34 Shumway N E. Personal communication. Unpubl 1972
 - 35 Eli R J and Zabriskie J H. Heart reactive antibody. A monitor of cardiac rejection. *Transplantation Proc* 3:905 1971
 - 36 Benzing G III, Spitzer R E, Bove K E, Schreiber J T and Helmsworth J A. Detection and treatment of canine cardiac rejection. *Transplantation* 14:35 1972
 - 37 Tenenbaum J I, Vasko J S and St Pierre R L. Canine heart allograft rejection and spontaneous in vitro transformation of peripheral blood lymphocytes. *Am J Med Sci* 258:59 1969
 - 38 Kahn D R, Carr E A, Dufek J H, Kursh M N, Gago O, Moores W Y, Oberman H A, Carroll M and Sloan H. Diagnosis of chronic rejection after cardiac transplantation in humans. *Transplantation Proc* 3:380 1971
 - 39 Deodhar S D, Konomi K, Nakamoto S and Kuruwita K C. Clinical experience with antilymphocyte globulin (ALG) in renal transplantation. *Transplantation Proc* 3:758 1971
 - 40 Deodhar S D, Kuklin A G, Vidt D G, Robertson A L and Hazard J B. Development of reticulum cell sarcoma at the site of antilymphocyte globulin injection in a patient with renal transplant. *N Engl J Med* 280:1104 1969
 - 41 Moore T C. In discussion of Stinson and associates. *Reference* 32
 - 42 Graham W H, Childs J W, DeGiorgi L S, Weymouth R J, Seibel H R and Lower M R. The effect of local graft irradiation on rejection of canine cardiac allografts. *J Thorac Cardiovasc Surg* 60:730 1970
 - 43 Wolf J S, McGavic J D and Hume D M. Inhibition of the effector mechanism of transplant immunity by local graft irradiation. *Surg Gynecol Obstet* 128:584 1969
 - 44 Kauffman H M Jr, Cleveland R J, Robertson G E, Graham W H and Hume D M. Inhibition of the afferent arc of the immune response to renal homografts by local graft irradiation. *Surg Gynecol Obstet* 123:1052 1966
 - 45 Miller J F A P. The thymus. Yesterday, today, and tomorrow. *Lancet* 2:1299 1967
 - 46 Snell G D. Immunologic enhancement. *Surg Gynecol Obstet* 130:1109 1970
 - 47 Nossal G J V. Summary of the Third International Congress of Transplantation Society. A personal approach. *Transplantation Proc* 3:1967 1971
 - 48 Griep R B, Wexler L, Stinson E B, Dong E Jr and Shumway N E. Coronary arteriography following cardiac transplantation. *JAMA* 221:147 1972
 - 49 Kahn D R, Carr E A Jr, Oberman H A, Kursh M M, Dufek J H, Moores W Y, Carroll M, Gago O and Sloan H. Effect of anticoagulants on the transplanted heart. *J Thorac Cardiovasc Surg* 60:616 1970
 - 50 Kincaid Smith P. Modification of the vascular lesions of rejection in cadaveric renal allografts by Dipyridamole and anticoagulants. *Lancet* 2:920 1969
 - 51 Kosek J C, Bieber C P and Lower R R.

or insurance to the patient. It is unlikely that such conditions can be met without adequate federal support.

Is the procedure worth this tremendous burden, particularly the financial one, at a time when most of our health institutions are in deep financial troubles? We elect to let Sir Peter Medawar²⁶ answer this last question. "I sometimes think there is an almost willful misunderstanding of the benefits that transplantation may bring with it. Plaintive and despondent voices tell us that the effort expended upon these heroic surgical procedures is out of all proportion to the number that can benefit from it, that we are squandering our material and intellectual resources on these adventures at a time when 99 per cent of the world's population cries out for elementary medical care; that the number of patients who have carried grafts of kidneys for more than a few months is smaller than the number of people engaged in the research on transplantation."

This is an unjust criticism. All great advances in medicine must nowadays start with a great capital investment of time and manpower and energy. There was a time when the same criticism could have been leveled against research on antibiotics. I was working in Oxford when research on penicillin in Professor Florey's entire laboratory was brought to focus in the treatment of just one small boy. The point is that the heroic adventures of today are part of tomorrow's ordinary medical care. If we thwart or discourage research on transplantation today, we are deliberately willing away part of the medical heritage of the future.*

Human cardiac transplantation is no longer a technical surgical problem. A surgeon should not be permitted to perform this operation simply because he can and wants to do it. Should this be permitted, the operation would qualify for Shaw's definition of "murderous absurdity"²⁷ and Forsmann's initial evaluation would not be too far from the truth.

At present human cardiac transplantation should remain in the province of the

small number of groups with a profound commitment and with financial support.

REFERENCES

1. News media and personal communication with Lower R R Sept 1972
2. Chicago Sun Times Sunday July 9 1972
3. Lind VI 180 Quoted in Precope J. Medicine, magic and mythology. London 1954. William Heinemann Ltd p 160
4. Spivak C D. Jewish encyclopedia 8:409 1904
5. Winkler H. Ueber Propfbastarde und phänotypische chimären. Ber Dtsch Botanische Ges 1907
6. Carrel A. La technique opératoire des anastomoses vasculaires et la transplantation des viscères. Lyon Med 98:859 1907
7. Carrel A. Latent life of arteries. J Exp Med 12:460 1910
8. Kolff W J. Removing limiting factors for total cardiac replacement. Transplantation Proc 3:1449 1971
9. Mann F C, Priestley J T, Marcowitz J and Yates W M. Transplantation on the intact mammalian heart. Arch Surg 261:19 1933
10. Carrel A and Guthrie C C. The transplantation of veins and organs. Am Med 11:1101 1905
11. Hardy J D, Chavez C M, Kurrus F D, Neely W A, Lishin S, Turner M D, Favian I W and Labecki T D. Heart transplantation in man. Developmental studies and report of a case. JAMA 188:1132 1964
12. Barnard C N. A human cardiac transplant. S Afr Med J 41:1271 1967
13. Edwards J E and Lorschmann W, quoted by Dubost C. Scientific and ethical problems in organ transplantation. Honored guest address presented at the Fifth Annual Meeting of the Society of Thoracic Surgeons, San Diego Calif Jan 27-29 1969. Published in Ann Thorac Surg 8:195 1969
14. American College of Surgeons. National Institutes of Health. Organ Transplant Registry. First Scientific Report. JAMA 217:1520 1971
15. Lower R R, Steffer R C and Shumway N E. Homovital transplantation of the heart. J Thorac Cardiovasc Surg 41:196 1961
16. Dong L Jr. Stanford University Medical Center News Bureau. Stanford Calif May 24 1972
17. Patterson S W, Piper H and Starling E H. The regulation of the heart beat. J Physiol (Lond) 48:465 1914
18. Grasser H W and Meek W J. Study of the mechanisms by which muscular exercise produces acceleration of the heart. Am J Physiol 34:48 1914
19. Samrian A. Muscular work in dogs submitted to different conditions of cardiac and splanchnic innervations. J Physiol (Lond) 83:313 1935
20. Cooper T, Gilbert J W, Bloodwell R D and Crout J R. Chronic extrinsic cardiac denervation

*Medawar. Sir Peter Medawar's views on transplantation. Br Med J 1:373 1968. Quoted with permission.

Electrocardiogram and vectorcardiogram in ventricular inversion (corrected transposition)

Benjamin E Victorica MD*

B Lynn Miller MD**

Ira H Gessner MD***

Gainesville Fla

Inversion of the ventricles with transposition of the great arteries (congenital corrected transposition of the great arteries) is a well defined cardiac malformation in which there is a reversal of the left right relationship of the ventricles with anatomic transposition of the aorta and pulmonary artery.^{1,2} The posterior pulmonary artery receives systemic venous blood from a morphologic left ventricle which functions as the venous ventricle while the anterior aorta receives pulmonary venous blood from a morphologic right ventricle that serves as the arterial ventricle. Thus the circulatory pathway is physiologically correct. However this entity rarely occurs without associated cardiac malformations. Ventricular septal defects, pulmonic stenosis, or an Ebstein like anomaly involving the atrioventricular valve of the morphologic right ventricle are commonly present.³⁻⁶

Several investigators^{3,5,10} have described the electrocardiographic (ECG) and vectorcardiographic (VCG) features of corrected

transposition but few cases have been correlated with the hemodynamic data. Furthermore some authors have included varieties of single ventricle with inversion in the analysis.¹⁷ The purpose of this report is to present the ECG VCG patterns observed with different hemodynamic states in a series of patients with ventricular inversion and two functioning ventricles. All had associated intracardiac malformations of varying severity. We have found that the ECG and VCG are reliable indicators of ventricular overloading just as they are in similar defects in the non inverted heart.

Materials and methods

We have reviewed nine children with ventricular inversion who were evaluated at the University of Florida Teaching Hospital and met our strict diagnostic criteria. Only patients with apex on the left were included in the series. In all cases the diagnosis of ventricular inversion with two functioning ventricles was based on cardiac catheterization and selective ventricular

*In the Division of Pediatric Cardiology, Department of Pediatrics, University of Florida College of Medicine, Gainesville, Fla.

Supported in part by the Departmental Physiology Training Grant, National Institutes of Health, TI HD-0054 and by Grants No. 71 AG-23, 71 AG-10 and No. 0-ARF 11 from the Florida Heart Association. Dr. Gessner is the Chief of the Cardiology Department, National Institutes of Health, HE 45142-04.

Received for publication, June 9, 1973.

Reprints request to: Benjamin E. Victorica, MD, Department of Pediatrics, University of Florida College of Medicine, Gainesville, Fla. 32601.

Assistant Professor of Pediatric Cardiology.

Assistant Professor of Pediatric Cardiology.

Chief Professor of Pediatric Cardiology, Head, Division of Pediatric Cardiology.

- Heart graft arteriosclerosis Transplantation Proc 3:512 1971
- 52 Beaumont J L Hyperlipidemia with circulating anti beta lipoprotein auto-antibody in man Auto-immune hyperlipidemia its possible role in atherosclerosis Progr Biochem Pharmacol 11:10 1968
 - 53 Kondo Y Gridel I and Kindrowitz A Heart homotransplantation in puppies Long survival without immunosuppressive therapy Circulation 31 (Suppl 1):181 1965
 - 54 Lower R R Stofer R C Hurley E J Dong L Jr Cohn R B and Shumway N L Successful homotransplantation of the canine heart after anoxic preservation for seven hours Am J Surg 104:302 1962
 - 55 Manax W J Iargader I and Lillehei R C Whole canine organ preservation JAMA 196:121 1966
 - 56 Lyons G W Dietzman R H and Lillehei R C On the mechanism of preservation with hypothermia and hyperbaric oxygen Trans Am Soc Artif Intern Organs 12:236 1966
 - 57 Kountz W B Revival of human hearts Ann Intern Med 10:330 1936
 - 58 Carrel A and Lindbergh C A The culture of whole organs Science 81:621 1935
 - 59 Dietrich E H Liddicoat J L Schwartz A Sordahl I A Brooks D K and DeBakey M I Preservation of the human heart Lx habit and brochure produced by medical illustration and audiovisual education Baylor College of Medicine 1970
 - 60 Belzer G O Ashby D S and Dunphy J E Twenty four hour and 72 hour preservation of canine kidneys Lancet 2:536 1967
 - 61 Korow A M Jr Webb W R and Stapp J E Preservation of hearts by freezing Arch Surg 91:572 1965
 - 62 Theodorides T Webb W R Nakae S and Sugg W L Prolonged survival of the anoxic lung with metabolic inhibitors in normal thermic and hypothermic conditions Ann Thorac Surg 5:411 1968
 - 63 Webb W R Jones F V and Sugg W L Magnesium and transmembrane potentials Fed Proc 27:579 1968
 - 64 Angell W W and Shumway N E Resuscitation storage of the cadaver heart transplant, Surg Forum 17:224 1966
 - 65 Wells P H Phalakornkul S Ramsey H W and Wheat M W Jr Cardiac function after prolonged storage in an intermediate biologic host J Thorac Cardiovasc Surg 62:869 1971
 - 65a Gertz A quoted by Ayd F J Jr What is death? Presented at the Second National Congress on Medical Ethics sponsored by the Judicial Council of the American Medical Association October 5-6 1968 Chicago Ill
 - 66 Sadler A M Jr Sadler M L and Stason C D JAMA 206:2501 1968 Recent development in the legal aspects of transplantation in the US Transplantation Proc 3:793 1971 Kansas State Ann Ch 378 (H B 1961) 1970
 - 67 Bernard C Quoted from Status of Transplantation 1968 A report by the Surgery Training Committee of the National Institute of General Medical Sciences National Institutes of Health page 51
 - 68 Medawar P Sir Peter Medawar's views on transplantation Br Med J 1:1373 1968
 - 69 Shaw G B Preface on doctors-The doctors dilemma New York 1948 Dodd Mead & Company Inc

Electrocardiogram and vectorcardiogram in ventricular inversion (corrected transposition)

Benjamin E Victorica MD*
B Lynn Miller MD**
Ira H Gessner MD***
Gainesville Fla

Inversion of the ventricles with transposition of the great arteries (congenital corrected transposition of the great arteries) is a well defined cardiac malformation in which there is a reversal of the left right relationship of the ventricles with anatomic transposition of the aorta and pulmonary artery.^{1,2} The posterior pulmonary artery receives systemic venous blood from a morphologic left ventricle which functions as the venous ventricle while the anterior aorta receives pulmonary venous blood from a morphologic right ventricle that serves as the arterial ventricle. Thus the circulatory pathway is physiologically correct. However this entity rarely occurs without associated cardiac malformations. Ventricular septal defects, pulmonic stenosis, or an Ebstein like anomaly involving the atrioventricular valve of the morphologic right ventricle are commonly present.^{3,4}

Several investigators⁵⁻¹⁰ have described the electrocardiographic (ECG) and vectorcardiographic (VCG) features of corrected

transposition but few cases have been correlated with the hemodynamic data. Furthermore some authors have included varieties of single ventricle with inversion in the analysis.¹¹ The purpose of this report is to present the ECG/VCG patterns observed with different hemodynamic states in a series of patients with ventricular inversion and two functioning ventricles. All had associated intracardiac malformations of varying severity. We have found that the ECG and VCG are reliable indicators of ventricular overloading just as they are in similar defects in the non inverted heart.

Materials and methods

We have reviewed nine children with ventricular inversion who were evaluated at the University of Florida Teaching Hospital and met our strict diagnostic criteria. Only patients with apex on the left were included in the series. In all cases the diagnosis of ventricular inversion with two functioning ventricles was based on cardiac catheterization and selective ventricular

From the Department of Pediatrics, Cardiology Department, University of Florida College of Medicine, Gainesville, Fla.
Supported in part by the Department of Physiology, University of Florida, Gainesville, Fla. and by the Department of Pediatrics, University of Florida, Gainesville, Fla.
Reprints: Dr. B. E. Victorica, MD, Department of Pediatrics, University of Florida College of Medicine, Gainesville, Fla. 32601.
*Assistant Professor of Pediatrics, Cardiology.
**Associate Professor of Pediatrics, Cardiology.
***Professor of Pediatrics, Cardiology, Department of Pediatrics, University of Florida College of Medicine, Gainesville, Fla.

Table I Clinical findings—Ventricular inversion with two functioning ventricles

| | Case no. | Age at catheterization | Sex | Symptoms signs | Age murmur discovered | Method of diagnosis | Associated defects | Surgery |
|-----------|----------|------------------------|-----|--------------------------------------|-----------------------|---------------------|--|-----------------------------------|
| Group I | | | | | | | | |
| | 1 | 3-6/12 years | F* | None | At birth | C-A | Subvalvar pulmonic stenosis | None |
| | 2 | 8-7/12 years | | None | | | | |
| | 3 | 4-7/12 years | M | None | 8 weeks | C-A | Ventricular septal defect, pulmonic valvar stenosis, left A V valve incompetence | None |
| Group II | | | | | | | | |
| | 3 | 13-6/12 years | M | None | 11 years | C-A | Pulmonic valvar stenosis | None |
| | 4 | 3 months | M | Jaundice Congestive heart failure | At birth | C-A V | Atrial septal defect azygos continuation of the inferior vena cava biliary atresia | None |
| | 5 | 1 year | F | None | 11 months | C-A | Ventricular septal defect pulmonic valvar stenosis | None |
| | 6 | 1-6/12 years | F | Cyanosis | At birth | C-A | Ventricular septal defect pulmonic valvar stenosis, right aortic arch | None |
| Group III | | | | | | | | |
| | 7 | 6 weeks | M | Congestive heart failure | 3 weeks | C-A-S | Ventricular septal defect | Pulmonary artery banding |
| | 8 | 3 years | F | None | 2 years | C-A | Ventricular septal defect pulmonic stenosis | None |
| | 9 | 13-8/12 years | M | Chronic congestive heart failure | 10 days | C-A-S | Ventricular septal defects (membranous & muscular) coarctation pulmonary arteries, pulmonic valve incompetence | Closure—ventricular septal defect |

Abbreviations: F = female; N = necropsy; M = male; S = surgery; C-A = catheterization and angiocardiography; A-V = atrioventricular.

angiocardiography with clear demonstration of the internal anatomic characteristics of the ventricles. Diagnosis on position of the great arteries alone is not acceptable to us. All patients had situs solitus thus by definition excluding all patients with either proved or suspected asplenia or polysplenia syndromes. Basic clinical information and description of the associated defects is given in Table I.

The ECG VCG tracings used for analysis were those obtained at the time of admission for cardiac catheterization. Electrocardiograms were recorded on direct writing units and included the classical six standard and augmented limb leads and the eight precordial leads (V_1R to V_4). Tracings were analyzed for rhythm, mean axes and QRS-T contour. Vectorcardiograms were obtained with a Sanborn Visoscope Model

569 A using the Crishman cube system. Photographs of the oculo-scope patterns were made on a Hewlett Packard 196 A camera in the frontal (F), right sagittal (S) and horizontal (H) planes. The electron beam was interrupted every 2.5 msec with the blunt end of the dots indicating the direction of inscription. The QRS loops were analyzed for the direction of inscription and the orientation of initial and terminal 0.01 second instantaneous and maximal QRS forces.

Classification. Catheterization data presented in Table II provide a hemodynamic basis for dividing the series into three groups.

Group I consists of three patients with mild pulmonic stenosis in whom the systolic pressure in the venous ventricle was 50 per cent of systemic or less. Case No. 2 and 3

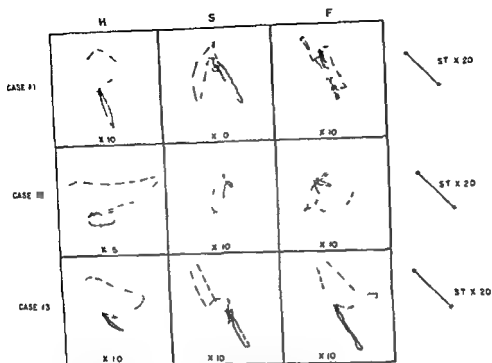


Fig. 1. Human cube vectorcardiograms of all three cases in Group I. Note the completely posterior counter-clockwise horizontal QRS loop in each case. Key: ST = 1.4 mm; \angle = standard and degree of attenuation; a row = direction of inscription; H = horizontal; S = right sagittal; F = frontal.

had a small ventricular septal defect with a pulmonary to systemic flow ratio (Q_p/Q_s) of 1.2:1 and moderate left atrioventricular valve incompetence. Case No. 1 is considered only in Group I although as noted in Table II and in the discussion below this patient progressed to meet the criteria for Group II at the time of a second catheterization.

Group II consists of three patients in whom the major hemodynamic effect was either volume or pressure overload of the venous ventricle. Case No. 4 had an atrial septal defect with a pulmonary to systemic flow ratio of 4:1. Cases No. 5 and 6 had ventricular septal defects associated with significant pulmonic valvular stenosis resulting in systemic pressures in the venous ventricle and net right to left shunts (Q_p/Q_s 0.93:1 and 0.6:1 respectively).

Group III consists of three patients with biventricular overload due to the presence of large ventricular septal defects. All had venous ventricular peak systolic pressure near systemic levels with left to right shunts ranging from 2.1:1 to 2.8:1. In

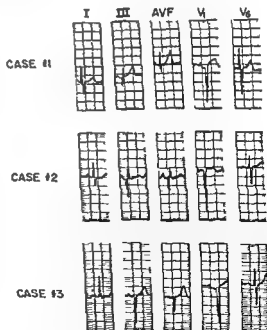


Fig. 2. Selected ECG leads from the three cases in Group I. Note the QS patterns in V_1 representing the posterior displacement of the QRS force in each case and the consistent q wave in Lead III and aVF.

Table II Catheterization data—Ventricular inversion with two functioning ventricles

| Case no | Pressures (mm Hg) | | | | |
|------------------|-------------------|---------------|-----------|----------------|--------|
| | V1* | P1 | 11 | 1a | 11/11a |
| Group I | | | | | |
| 1 | 50/2 5 | 17/8 (9)‡ | 100/0-5 | | 0 50 |
| 2 | 59/0-3 | 31/12 | 98/4 5 | 98/58 (80) | 0 40 |
| 3 | 48/0 6 | 30/10 (19) | 108/2 12 | 108/10 (90) | 0 44 |
| Group II | | | | | |
| 4 | 58/11 19 | | 110/18 32 | 115/35 | 0 53 |
| 5 | 81/0-4 | | 85/0-5 | 87/48 (58) | 0 99 |
| 6 | 90/2 8 | | 90/2 10 | 90/45 (68) | 1 0 |
| 1† | 65 (90)‡/3 6 | 15/9 (13) | 88/3 7 | 88/60 | 0 44 |
| Group III | | | | | |
| 7 | 80/0-5 | 83/32 (45) | | 98/50 (68) | 0 82 |
| 8 | 90/5 10 | 27/15 | | 116/58 (10) | 0 8 |
| 9 | 87/8 14 | 85/36 | 107/7 20 | 104/10 | 0 81 |

Abbreviations: VV = venous ventricle; 11 = pulmonary artery; 11 = arterial ventricle; 1a = aorta; V1/11 or 1a = peak systolic pressure.

† = second catheterization.

‡ = after angiocardioradiography.

§ Parentheses denote mean pressures.

NOTE: Pulmonary blood flow in Cases No. 5 and 6 was estimated assuming a pulmonary venous saturation of 97 per cent. Indicator dilution method.

In addition, Case No. 6 demonstrated pulmonic valve stenosis and Case No. 9 was found to have bilateral coarctations of the pulmonary arteries and pulmonic valve incompetence.

Results

Pertinent electrocardiographic and vectorcardiographic information is given in Tables III and IV.

Group I (Cases No. 1 to 3) The initial (0.01 sec) QRS vector is oriented directly leftward or leftward posteriorly and superiorly (Fig. 1). The QRS loop is completely posterior in all predominantly inferior in two and superior in one. The terminal QRS forces are directed rightward and posteriorly in all superiorly in two and inferiorly in one. The QRS loops in the H plane are very similar in the three cases being open loops with counterclockwise

(CCW) inscription. Frontal plane QRS inscription ranges from predominantly CCW in one to purely clockwise (CW) in another.

The electrocardiograms (Fig. 2) demonstrate a mean QRS axis in the frontal plane of +25 degrees in two cases and -30 degrees in one. Initial q waves present in Leads III and aVF in all three are deeper in Lead III in each instance. The QRS morphology in the precordial leads is uniform in all cases reflecting the posterior displacement of the forces. A consistent QS pattern is present in V1 while in V6 there are no initial q waves but rather RS complexes.

Group II (Cases No. 4 to 6) Vectorcardiograms (Fig. 3) reveal initial QRS forces directed leftward and superiorly in every case. In two the 0.01 sec vector is posterior and in one directly leftward at 0

1 May 86
Number 6

| Case | Oxygen saturation (%) | | | | | Qp/Qs | Selective angiocardiology |
|------|-----------------------|----|----|----|----|-------|--|
| | RA | IV | PA | AV | Ao | | |
| 81 | | | 67 | 98 | | 1.1 | None |
| 82 | 64 | | 10 | 98 | | 1.2 | Venous ventricle arterial ventricle |
| 61 | | 90 | 81 | | 97 | 1.1 | Venous ventricle |
| 61 | 54 | 57 | | | 98 | 4.5 | Venous ventricle arterial ventricle |
| 61 | 62 | 62 | | | 91 | 0.93 | Venous ventricle |
| 72 | | | 74 | | 83 | 0.6 | Venous ventricle |
| 81 | | | 84 | | 98 | 1.1 | Venous ventricle |
| 81 | | | 78 | | 95 | 2.1 | Venous ventricle left atrium |
| 81 | 80 | | 91 | | 93 | 2.3 | Venous ventricle |
| 81 | | | | | 97 | 2.8 | Arterial ventricle pulmonary artery |

VC = ventricular catheter RA = right atrium Qp = pulmonary blood flow Qs = systemic blood flow

Method: modified right-to-left shunt Case No. 8

degrees. The QRS loops curiously have a similar configuration in the H and F planes with predominantly CW inscription and the major portion of the loop directed rightward anteriorly and inferiorly in every case. The terminal QRS forces are uniformly rightward anterior and inferior.

The mean frontal QRS axis on the ECC (Fig. 4) is +130 degrees in all three cases and all have q waves in Leads III and aV_r. Unlike Group I a qR pattern in the right precordial leads is the rule. There are no initial q waves in V₁, all demonstrating RS complexes. These ECG/VCG findings are considered to be indicative of hypertrophy of the venous ventricle.

Group III (Cases No. 7 to 9) In the VCG (Fig. 5) the initial (0.01 sec) QRS forces are all leftward, two being superior and one inferior. In contrast with the two previous groups, the initial QRS forces are anteriorly

directed in every case. In Case No. 7 following the initial QRS vector the H plane QRS loop continues leftward but then the loop reverses and the major portion is rightward and anterior with a CW inscription. The vectorcardiograms in Cases No. 8 and 9 show a different pattern with large anterior and posterior QRS forces and CCW inscription in the H plane. The F plane in Case No. 8 shows CCW inscription of the QRS while Cases No. 7 and 9 are CW.

In the electrocardiograms (Fig. 6) the mean QRS axis in the F plane ranges between +50 degrees and +100 degrees. An initial q wave is present in Lead III in all but no q waves are present in the precordial leads. Leads V₁ and V₂ show RS complexes of varying magnitude but Lead V₁ in Case No. 7 demonstrates an rSR pattern. Large biphasic mid precordial QRS complexes

Table II Catheterization data—Ventricular immersion with two functioning ventricles

| Case no | Pressures (mm Hg) | | | | |
|------------------|-------------------|----------------|-----------|----------------|----------------------------------|
| | VV* | PA | LV | Ao | V ₁ /V ₄ † |
| Group I | | | | | |
| 1 | 50/2.5 | 17/8 (9)‡ | 100/0.5 | | 0.50 |
| 2 | 39/0.3 | 31/12 | 98/4.5 | 98/58 (80) | 0.40 |
| 3 | 48/0.6 | 30/10 (19) | 108/2.12 | 108/10 (90) | 0.44 |
| Group II | | | | | |
| 4 | 58/11.19 | | 110/18.32 | 115/35 | 0.53 |
| 5 | 84/0.4 | | 85/0.5 | 82/48 (58) | 0.99 |
| 6 | 90/2.8 | | 90/2.10 | 90/45 (68) | 1.0 |
| 1† | 65 (90)‡/3.6 | 15/8 (13) | 88/3.7 | 88/60 | 0.14 |
| Group III | | | | | |
| 7 | 80/0.5 | 81/3.7 (45) | | 98/50 (68) | 0.87 |
| 8 | 90/5.10 | 27/15 | | 116/58 (70) | 0.18 |
| 9 | 87/8.14 | 85/36 | 107/7.20 | 104/70 | 0.81 |

*Abbreviations: VV = venous ventricle; PA = pulmonary artery; LV = arterial ventricle; Ao = aorta; V₁/V₄ or A_o = peak systolic pressure.

† = second catheterization.

‡ = after angiocardiology.

§ Parentheses denote mean pressures.

NOTE: Pulmonary blood flow in Cases No. 5 and 6 was estimated assuming a pulmonary venous saturation at 97 per cent. Indicator dilution method.

In addition, Case No. 8 demonstrated pulmonic valve stenosis and Case No. 9 was found to have bilateral coarctations of the pulmonary arteries and pulmonic valve incompetence.

Results

Pertinent electrocardiographic and vectorcardiographic information is given in Tables III and IV.

Group I (Cases No. 1 to 3) The initial (0.01 sec) QRS vector is oriented directly leftward, or leftward posteriorly and superiorly (Fig. 1). The QRS loop is completely posterior in all, predominantly inferior in two and superior in one. The terminal QRS forces are directed rightward and posteriorly in all, superiorly in two and inferiorly in one. The QRS loops in the II plane are very similar in the three cases, being open loops with counterclockwise

(CCW) inscription. Frontal plane QRS inscription ranges from predominantly CCW in one to purely clockwise (CW) in another.

The electrocardiograms (Fig. 2) demonstrate a mean QRS axis in the frontal plane of +25 degrees in two cases and -50 degrees in one. Initial q waves present in Leads III and aV_F in all three are deeper in Lead III in each instance. The QRS morphology in the precordial leads is uniform in all cases, reflecting the posterior displacement of the forces. A consistent QS pattern is present in V₁ while in V₄ there are no initial q waves but rather RS complexes.

Group II (Cases No. 4 to 6) Vectorcardiograms (Fig. 3) reveal initial QRS forces directed leftward and superiorly in every case. In two the 0.01 sec vector is posterior and in one directly leftward at 0

Table IV Vectorcardiographic findings—Ventricular inversion with two functioning ventricles

| Horizontal plane | | | | | | Right sagittal plane | Frontal plane | | | | |
|------------------|-----------------|------|-----------------|------|-----|----------------------|-----------------|------|-----------------|--------|-----|
| QRS loop | | | | | | T loop | QRS loop | | | T loop | |
| Case no. | Init (0.01 sec) | Max | Term (0.01 sec) | Insc | Max | Insc | Init (0.01 sec) | Max | Term (0.01 sec) | Insc | Max |
| Group I | | | | | | | | | | | |
| 1 | +35 | +90 | +130 | CCW | -0 | CW | -40 | +50 | -140 | CW | +10 |
| | +15 | +30 | +150 | CCW | -30 | CCW | -40 | +170 | +150 | CW | +15 |
| 3 | 0 | +5 | +1.5 | CCW | -45 | CW | -10 | -110 | -170 | CCW | +60 |
| Group II | | | | | | | | | | | |
| 4 | +70 | -160 | -150 | CW | | CCW | -45 | +160 | +160 | CW | +80 |
| 5 | -10 | -150 | -130 | CW | -10 | = | -10 | +165 | +160 | = | +5 |
| | | | | (=) | | (CCW) | | | | (CW) | |
| 6 | -5 | -170 | -130 | CW | | CCW | -10 | +60 | +150 | CW | +60 |
| 1† | +35 | -150 | -150 | CW | -40 | CCW | -60 | +45 | +160 | CW | +0 |
| Group III | | | | | | | | | | | |
| 7 | -60 | -10 | -170 | CW | | CCW | -40 | +100 | +130 | CW | +80 |
| 8 | -65 | +65 | +80 | CCW | -80 | CW | +30 | +40 | +35 | CCW | +90 |
| 9 | -65 | +120 | -140 | CCW | -60 | = | -30 | +120 | +145 | CW | +60 |
| | | | | | | (CW) | | | | | |

Abb: 1 CW = clockwise CCW = counter clockwise = = complex O = in the horizontal plane & it is considered as negative I at Max T m = 1st max max. and 2nd max max. QRS for I sec = direction of inscription of the QRS loop
† = at second time

are present in all three cases. These ECG VCG findings are felt to represent biventricular hypertrophy.

Discussion

Leftward and superior orientation of the initial QRS forces in inversion of the ventricles is a consistent and well-documented phenomenon (Fig. 7). This reversal of septal depolarization is attributed to inversion of the conduction pathways (atrioventricular node, H₁ bundle and bundle branches) clearly shown by the work of Lev, Licata and May.¹⁹ The resultant ECG manifestations are the presence of qR patterns in Leads III and aV_F and reversal of the precordial Q wave pattern as first described by Anderson and co-workers.⁵ Little emphasis, however, has been placed on the study of the orientation and configuration of the rest of the QRS forces, particularly as they relate to the presence or absence of associated cardiac malformations and their hemodynamic consequences (ventricular

overload). By limiting our analysis to unequivocal cases of ventricular inversion with situs solitus cardiac apex on the left and two functioning ventricles we have confirmed the presence of consistent ECG VCG patterns.

In cases of inversion with minor associated anomalies resulting in low venous ventricular pressure (our Group I) the ECG VCG shows a pattern which may be considered normal for this cardiac condition.^{7, 12, 14} In these patients the VCG shows maximal and terminal QRS forces oriented posteriorly and the QRS loops in the H plane are open and wide with CCW inscription. The few published vector cardiograms whether recorded by the Frank²⁰ or Grishman²¹ system show similar configurations.^{1, 14} The ECG pattern of QS complexes in the right precordial leads (V₁R to V₁) seen in our patients has been described in isolated cases in the literature.^{8, 10, 22} Occasional cases with small r deep S in Lead V₁ have been seen.¹⁴

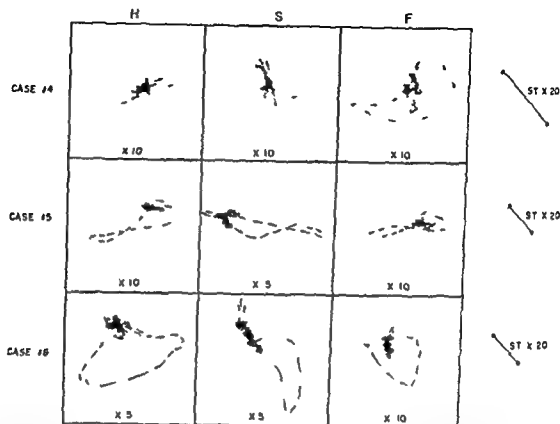


Fig. 3. Cube vectorcardiograms in all three cases of Group II. Note that while the mutual QRS forces are oriented leftward either at 0 degrees or more posteriorly, the major QRS forces are anterior and rightward in each case. Key same as Fig. 1.

Table III. *Electrocardiographic findings—1 ventricular inversion with two functioning ventricles*

| Case no | Rhythm | P | Mean axis (degrees) | | PR (sec) | Q/R/S Pattern (mm) | | | | | T II area | |
|-----------|-----------------------|------|---------------------|------|-----------|--------------------|--------|---------|---------|-----|-----------|----|
| | | | QRS | T | | III | aVF | I | II | III | I | II |
| Group I | 1 *NSR | +60 | +25 | +10 | 0.12 | 5/5/0 | 2/1/4 | 10/0/10 | 0/11/13 | + | + | + |
| | 2 Variable AV Block | +65 | +25 | +40 | 0.16 | 10/6/0 | 6/6/0 | 20/0/0 | 0/1/3 | + | + | + |
| | 3 NSR 1 AV Block | +60 | -30 | +0 | 0.0 | 10/0/7 | 6/0/5 | 10/0/0 | 0/10/10 | + | + | + |
| Group II | 4 2 AV Block | +50 | +130 | -50 | 0.11 | 13/18/0 | 7/10/0 | 3/5/0 | 0/7/16 | + | + | + |
| | 5 Wandering pacemaker | - | +130 | +10 | 0.10-0.11 | 4/11/0 | 2/5/0 | 2/1/1 | 0/6/10 | + | + | + |
| | 6 NSR | +60 | +130 | +60 | 0.11 | 7/13/0 | 5/15/0 | 4/13/0 | 0/8/10 | + | + | + |
| Group III | 1† NSR | +60 | +70 | +80 | 0.12 | 5/11/1 | 2/9/3 | 10/16/0 | 0/10/15 | + | + | + |
| | 7 NSR | +60 | +110 | +60 | 0.10 | 5/17/0 | 6/28/1 | 0/1/1/6 | 0/15/14 | + | + | + |
| | 8 NSR | +10 | +30 | +90 | 0.11 | 2/7/0 | 0/8/0 | 0/10/4 | 0/10/3 | + | + | + |
| | 9‡ NSR | | | | | | | | | | | |
| | 1 AV Block | +100 | +60 | +110 | 0.06 | 4/3/5/9 | 3/8/0 | 0/10/5 | 0/1/19 | + | + | + |

*Abbreviations: AV = atrioventricular; NSR = normal sinus rhythm.

† = second catheterization.

‡ = complete AV block after surgery.

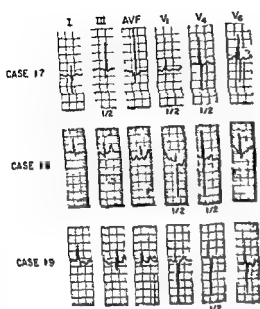


Fig 6 Selected ECG leads in all three cases of Group III. Note the absence of q waves both in the right and left precordial lead. Presence of a q in Lead III and large biphasic QRS complexes in V_1 . I_s standardization 10 mv = 5 mm

DISTRIBUTION OF INITIAL (0.01) QRS VECTORS

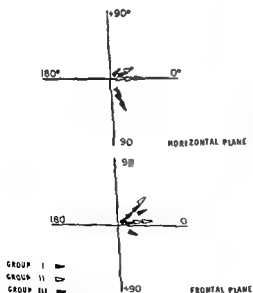


Fig 7 Composite of the initial (0.01 sec) QRS vector in the horizontal and frontal planes for all 9 cases of inversion of the ventricles with two functional ventricles. In the horizontal plane the anterior quadrants are considered negative while the posterior are positive.

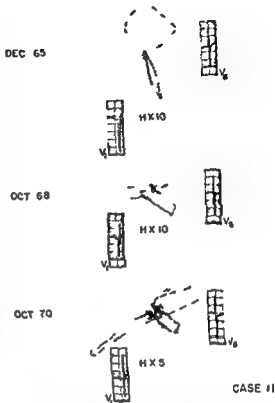


Fig 8 Serial H plane vectorcardiograms in Case 1 showing progression from loops characteristic of Group I (Dec 65) to Group II (Oct 70). Note that the QRS pattern in V_1 changes from a QS to a qR.

from that observed in double inlet left ventricle and inversion.²² Furthermore a qR pattern in V_1 may be present in some cases of severe right ventricular hypertrophy in a heart with non inversion of the ventricles. In such cases the initial forces in the frontal plane will generally help distinguish between the two.

Those cases with hemodynamic evidence of overload of both ventricles (our Group III) demonstrate ECG VCC patterns consistent with biventricular hypertrophy. In striking contrast with the two previous groups the initial 0.01 sec QRS forces while leftward are distinctly anterior. This finding could not be explained by the orientation of the ventricular septum as determined by angiocardiography. The ECC representation of anterior displacement of the initial QRS forces is absence of q waves in both right and left precordial leads. In Case No 7 the leftward and CCW early

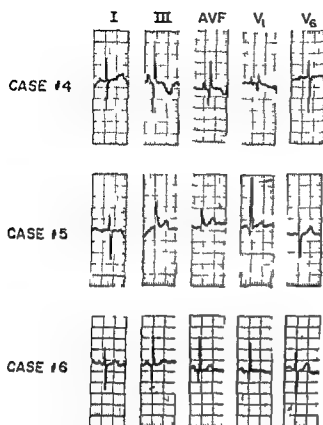


Fig 4 Selected ECG leads in all three cases of Group II. Note the right axis deviation, significant initial q waves in leads III and AVF, and constant QR pattern in V_1 .

In those cases with either volume or pressure overload of the venous ventricle (our Group II), the maximal and terminal QRS forces, in contrast with the previous group, have a rightward and anterior orientation as a result of the hypertrophy of the right sided morphologic left ventricle. The VCG in all three patients shows CW inscription of the QRS loop in the H plane. Ruttenberg and associates² have published a VCG which shows the same general configuration of the QRS from a child with similar hemodynamic findings.

The constant ECG feature observed in Group II is the well recognized qR pattern in V_1 generally described as a common finding in corrected transposition. This pattern was present in 60 per cent of the cases analyzed by Schiebler and associates.³ However, tall R waves in V_1 and predominant S waves in V_6 are signs of hypertrophy of the right sided ventricle. On review of the literature of corrected transposition with associated venous ventricle hypertension we found similar patterns.^{4, 6, 10, 2}

Present evidence suggests that the ECG VCG of corrected transposition with venous ventricular overload is indistinguishable

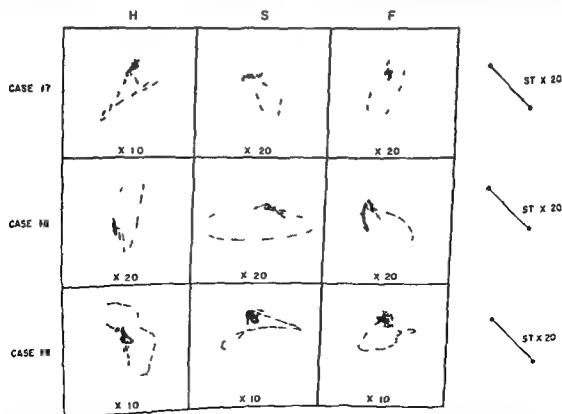


Fig 5 Cube vectorcardiograms in all three cases of Group III. Note that the initial QRS forces, while leftward and anterior in all. Cases No. 8 and 9 show large anterior and posterior QRS forces. Key same as Fig. 1.

Vol. 86
No. 6

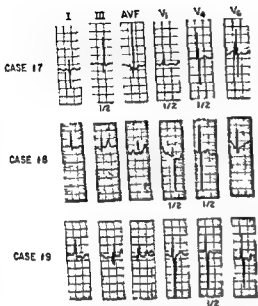


Fig 6 Selected ECG leads in all three cases of Group III. Note the absence of q waves both in the right and left precordial lead, presence of a q in Lead III and large biphasic QRS complexes in V_1 . $1\frac{1}{2}$ standardization 10 mv = 5 mm

DISTRIBUTION OF INITIAL 0.01 sec QRS VECTORS

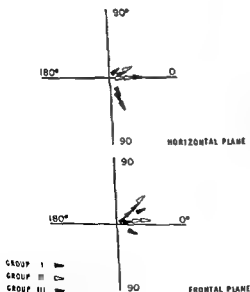


Fig 7 Composite of the initial (0.01 sec) QRS vector in the horizontal and frontal planes for all 9 cases of inversion of the ventricles with two functioning ventricles. In the horizontal plane the anterior-dorsal are considered negative while the posterior are positive.

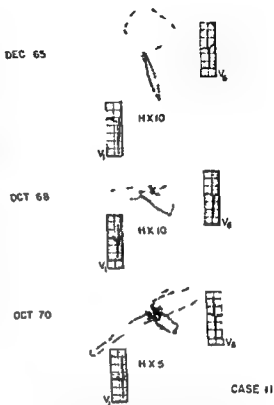


Fig 8 Serial H plane vectorcardiograms in Case No 1 showing progression from loops characteristic of Group I (Dec 65) to Group II (Oct 70). Note that the QRS pattern in V_1 changes from a QS to a qR.

from that observed in double inlet left ventricle and inversion.²² Furthermore, a qR pattern in V_1 may be present in some cases of severe right ventricular hypertrophy in a heart with non-inversion of the ventricles. In such cases the initial forces in the frontal plane will generally help distinguish between the two.

Those cases with hemodynamic evidence of overload of both ventricles (our Group III) demonstrate ECG/VCG patterns consistent with biventricular hypertrophy. In striking contrast with the two previous groups the initial 0.01 sec QRS forces while leftward are distinctly anterior. This finding could not be explained by the orientation of the ventricular septum as determined by angiocardiography. The ECC representation of anterior displacement of the initial QRS forces is absence of q waves in both right and left precordial leads. In Case No 7 the leftward and CCW early

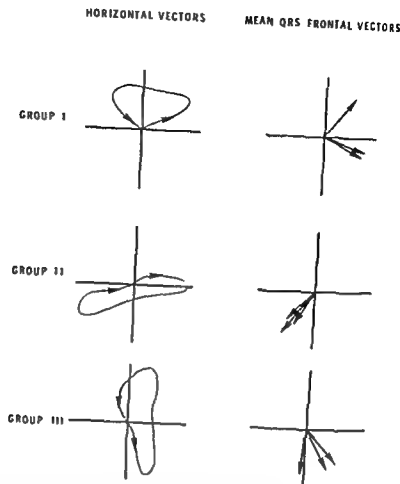


FIG 9 Derived Grijman system horizontal plane QRS loop and observed mean frontal QRS vectors in ventricular inversion with two functioning ventricles (corrected transposition). Group I minor associated defects. Group II venous ventricular overload. Group III biventricular overload.

QRS forces are followed by reversal of inscription to a larger rightward and anterior portion of the QRS loop. This represents biventricular hypertrophy, right greater than left in a patient who is only six weeks old. The horizontal QRS loops of cases No. 8 and 9 have a similar open and CCW inscription with large anterior and posterior forces which is not unlike a case shown by Ruttenberg and co-workers.²²

The ECG criteria for biventricular hypertrophy are large biphasic RS complexes in the mid precordial leads. Such prominent voltages are not present in any of the six cases in Groups I and II.

Case No. 1 presents a narrative further substantiating the impression that the ECG VCG described in the different groups illustrates varying physiologic states. Serial tracings in this patient (Fig. 8) show an evolution of the characteristics associated with Group I to Group II as the venous ventricular pressure increased due to progressive pulmonic stenosis.

Derived H plane QRS loops and plotted mean F plane QRS vectors (Fig. 9) demonstrate the features which best distinguish ventricular overload states in corrected transposition. In cases with minor associated anomalies the QRS loop is posterior with CCW inscription and there is borderline or frank left axis deviation. With venous ventricular overload the QRS loop is predominantly anterior with CW inscription and there is right axis deviation. Biventricular overload deviates the initial QRS forces anteriorly; the QRS loop may be leftward and CCW as shown or more rightward and anterior if the venous ventricle overload is predominant. The mean frontal QRS vector may be in normal range.

Summary

The vectorcardiographic (VCG) and electrocardiographic (ECG) features in nine cases of ventricular inversion with two functioning ventricles (congenital corrected transposition of the great arteries) and

1 of 1086
1 mb 6

associated cardiac malformations were correlated with their hemodynamic states. All three cases with low venous ventricular pressure showed a similar VCC pattern with a posterior and completely counter clockwise horizontal QRS loop. The ECC showed a leftward QRS axis. QS complexes in V_1 and RS complexes in V_4 . These findings are normal for ventricular inversion without major associated defects. Three patients with isolated pressure or volume overload of the venous ventricle showed large leftward and posterior initial QRS forces followed by a predominantly clockwise and rightward horizontal QRS loop. The ECC showed right axis deviation. QR complexes in V_1 and rS complexes in V_4 . These findings are considered to represent venous ventricular hypertrophy. Three patients with hemodynamic situations which would be expected to result in biventricular overload demonstrated varying degrees of anterior displacement of the horizontal QRS loop two with counter clockwise inscription and one predominantly clockwise. The initial QRS forces although leftward and superior as in the other groups were anterior in each case. The ECG demonstrated normal or rightward QRS axis, no precordial q waves and large biphasic RS complexes in the mid precordial leads. These findings are considered to represent biventricular hypertrophy. One patient had serial tracings which revealed progressive ECC/VCC changes due to increasing venous ventricular hypertension secondary to increasing pulmonary stenosis.

These findings indicate that the ECC/VCC accurately reflect the anatomy and hemodynamics in ventricular inversion.

REFERENCES

- 1 Von Rokkansky K F Die Defecte des Scheidewande des Herzens Vienna 1875 W B Baumüller p 83
- 2 Cardell H S Corrected transposition of the great vessels Br Heart J 18:186 1956
- 3 Schiebler G L Edward J E Burchell H B DuShane J W Ongley P A and Wood E H Congenital corrected transposition of the great vessels a study of 33 cases Pediatrics (Suppl) 27:851 1961
- 4 Helmholz H F Jr Daugherty G W and Edwards J E Cardiac clinics CMAA Congenital mitral insufficiency in association with corrected transposition of the great vessels Report of probable clinical case and review of six cases studied pathologically Mayo Clin Proc 31:82 1956
- 5 Anderson R C Lilliches C W and Lester R G Corrected transposition of the great vessels of the heart A review of 17 cases Pediatrics 20:676 1955
- 6 Fink B W Adams F H McCall R A et al Corrected transposition of the great vessels without significant associated defects Pediatrics 21:381 1958
- 7 Goodman A H and Kuzman W J Functionally corrected transposition of the great vessels without significant associated defects Am Heart J 61:311 1961
- 8 Beck W Schrire A Vexelpoel I et al Corrected transposition of the great vessels Br Heart J 23:497 1961
- 9 Bidawi H S Bidar M K Habib A A et al Corrected transposition of the great vessels Report of 2 cases Am Heart J 62:119 1961
- 10 Honey M The diagnosis of corrected transposition of the great vessels Br Heart J 20:113 1963
- 11 Berry W B Roberts W C Morrow A G et al Corrected transposition of the aorta and pulmonary trunk Clinical hemodynamic and pathologic findings Am J Med 36:135 1964
- 12 Rotem C E and Hultgren H A Corrected transposition of the great vessels without associated defects Am Heart J 70:305 1965
- 13 Gaultfield W H Jr Bostock B and Verliff J K Corrected transposition of the great vessels with isolated pulmonary stenosis The paradox of pulmonary stenosis with physical signs of pulmonary hypertension Am J Cardiol 19:285 1967
- 14 Cumming G R Congenital corrected transposition of the great vessels without associated intracardiac anomalies A clinical hemodynamic and angiographic study Am J Cardiol 10:605 1962
- 15 Paul M H Van Praagh S and Van Praagh R Corrected transposition of the great arteries in Watson H ed Pediatric cardiology St Louis 1968 The C V Mosby Company p 611
- 16 Ruttenberg H D Corrected transposition of the great vessels in Moss A J and Adams F H eds Heart disease in infants, children and adolescents Baltimore 1968 The Williams & Wilkins Company p 553
- 17 Friedberg D Z and Nadas A S Clinical profile of patients with congenital corrected transposition of the great arteries N Engl J Med 282:1053 1970
- 18 Kossman C E (Chairman) et al Report of the Committee on Electrocardiography American Heart Association Recommendations for standardization of leads and of specifications for instruments in electrocardiography and vectorcardiography Circulation 35:583 1967
- 19 Lev M Licata R H and May R C The conduction system in mixed levocardia with ventricular inversion (corrected transposition) Circulation 28:232 1963
- 20 Frank E An accurate clinically practical

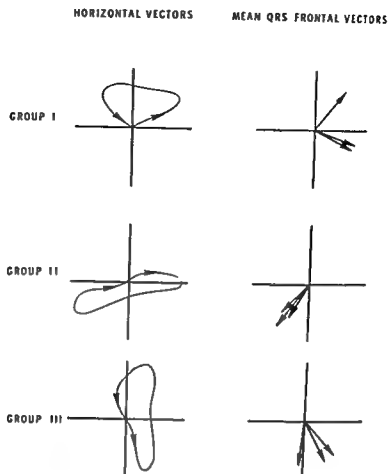


Fig 9 Derived Grishman system horizontal plane QRS loops and observed mean frontal QRS vectors in ventricular inversion with two functioning ventricles (corrected transposition). Group I minor associated defect. Group II venous ventricular overload. Group III biventricular overload.

QRS forces are followed by reversal of inscription to a larger rightward and anterior portion of the QRS loop. This represents biventricular hypertrophy, right greater than left in a patient who is only six weeks old. The horizontal QRS loops of cases No. 8 and 9 have a similar open and CCW inscription with large anterior and posterior forces which is not unlike a case shown by Ruttenberg and co-workers.²²

The ECG criteria for biventricular hypertrophy are large biphasic RS complexes in the mid precordial leads. Such prominent voltages are not present in any of the six cases in Groups I and II.

Case No. 1 presents a narrative further substantiating the impression that the ECG VCG described in the different groups illustrates varying physiologic states. Serial tracings in this patient (Fig. 8) show an evolution of the characteristics associated with Group I to Group II as the venous ventricular pressure increased due to progressive pulmonic stenosis.

Derived H plane QRS loops and plotted mean F plane QRS vectors (Fig. 9) demonstrate the features which best distinguish ventricular overload states in corrected transposition. In cases with minor associated anomalies the QRS loop is posterior with CCW inscription and there is borderline or frank left axis deviation. With venous ventricular overload the QRS loop is predominantly anterior with CW inscription and there is right axis deviation. Biventricular overload deviates the initial QRS forces anteriorly; the QRS loop may be leftward and CCW as shown or more rightward and anterior if the venous ventricle overload is predominant. The mean frontal QRS vector may be in normal range.

Summary

The vectorcardiographic (VCG) and electrocardiographic (ECG) features in nine cases of ventricular inversion with two functioning ventricles (congenital corrected transposition of the series) are:

Level of the base of the mitral valve

R W Broer Ph D

A H Krauss M D

G T Meester M D

Rotterdam The Netherlands

Mean left atrial pressure is a significant quantity in the assessment of left ventricular filling pressure and pulmonary venous pressure. These quantities in turn provide information concerning left ventricular performance and the development of pulmonary edema which are of real concern in the management of postoperative patients for heart and lung surgery. A number of guidelines have been presented in the literature for the determination of the zero pressure level for systemic veins for the patient supine.¹⁻⁴ However, the use of these guidelines for the left atrium and for patients in positions other than supine is in principle incorrect. Especially now that flow guided balloon catheters enable us to monitor pulmonary capillary wedge pressure in the Intensive Care Units a new investigation seems justified. In addition continuous monitoring of left atrial pressure after heart or lung surgery has shown significant changes in this parameter when patients were turned or tilted. This too might have been due to conventional methods for zero pressure levelling. Hence in order to determine whether these pressure changes were due to conventional methods for zero pressure

determination the vertical distance from the base of the mitral valve to a marker placed at mid sternum was determined radiologically. Furthermore we have investigated how well these distances correlated with thorax dimensions, patient height and weight.

Ten patients were chosen for this study and all were at least one month postoperative for replacement of the mitral valve (Starr Edwards ball valve). These patients were chosen since the mitral valve presented a clear unambiguous image on the x ray film and did not require any invasive procedures. Since a simple reference point was desirable all distances were measured with respect to the center of the sternum. The choice of the base of the mitral valve rather than some other point in the left atrium or left ventricle was dictated primarily by the patients available for this study. That is the base of the mitral valve represented the only clear unambiguous image on the x ray films to which distances could be easily measured. It can be argued however that this is not necessarily the most desirable point of reference with mid ventricle or mid atrium possibly a better choice.

Reprint requests to: Dr. R. W. Broer, The Thorax Center, Medische Faculteit, Rijkshospitaal, Postbus 1738, Rotterdam, The Netherlands.

This work was performed with partial support of the ZWO organization.

Received for publication January 9, 1973.

Reprint requests to: Dr. R. W. Broer, The Thorax Center, Medische Faculteit, Rijkshospitaal, Postbus 1738, Rotterdam, The Netherlands.

- system for spatial vectorcardiography. *Circulation* 13:737 1956
- 21 Grishman A, Borum R, and Jaffe H L. Spatial vectorcardiography. Technique for the simultaneous recording of the frontal, sagittal and horizontal projections. *Am Heart J* 41:483 1951
 - 22 Ruttenberg H D, Elliott L P, Anderson R C, et al. Congenital corrected transposition of the great vessels. Correlation of electrocardiograms and vectorcardiograms with associated cardiac malformations and hemodynamic states. *Am J Cardiol* 17:339 1966.
 - 23 Gessner I H, Elliott L P, Schiebler G L, et al. The vectorcardiogram in double inlet left ventricle with and without ventricular inversion. In Hoffman J, ed. *Vectorcardiography-2. Proceedings of the VI International Symposium on Vectorcardiography*. Amsterdam 1971. North Holland Publishing Company, p 624.

Table II Physical data on patients

| Patient no | Age (years) | Sex | Height (cm) | Weight (kg) | Surface area (M ²) | Sternum length (cm) | Thorax width (cm) | Thorax depth (cm) |
|------------|-------------|-----|-------------|-------------|--------------------------------|---------------------|-------------------|-------------------|
| 1 | 17 | F | 136 | 47.4 | 1.45 | 16 | 22 | 19 |
| 2 | 42 | F | 167 | 52.3 | 1.60 | 18 | 27 | 21 |
| 3 | 56 | F | 158 | 59.3 | 1.60 | 25 | 24.5 | 22 |
| 4 | 49 | F | 137 | 60 | 1.61 | 14.5 | 22 | 22.5 |
| 5 | 33 | F | 167 | 50 | 1.52 | 16 | 24 | 20 |
| 6 | 37 | M | 174 | 71.5 | 1.88 | 21 | 30 | 24 |
| 7 | 48 | M | 163 | 55.5 | 1.60 | 19 | 27 | 0 |
| 8 | 64 | M | 177 | 45 | 1.93 | 25 | 32 | 7 |
| 9 | 40 | F | 156 | 40.2 | 1.35 | 17 | 23.7 | 18 |
| 10 | 43 | F | 168 | 50.5 | 1.57 | 17.5 | 26.3 | 21 |

photo measurements the physiological distances for the projection used and (3) for each patient position use information from steps 1 and 2 from all camera positions to determine the three coordinates of the mitral valve with respect to the patient coordinate system.

Since all the patients had Starr Edwards ball valves some simplification in the methodology was possible. The diameter of the ball in the valve for each photo was used to determine the scale factor. This meant that only one calibrated photo need be taken to initially determine the actual ball diameter. For all subsequent pictures this diameter defined the scale factor with out any reference to source patient image distances.

Results

With the mid sternum marker as the reference point the data in Table I represent the location of the base of the mitral valve for all 10 patients in the 6 indicated positions. Within each box there are 4 numbers. The first three to the left in each box represent the x , y and $-z$ coordinates of the mitral valve with respect to the mid sternum marker. x represents the direction from the mid sternum marker to the feet, y the distance from the marker to the left arm and $-z$ the distance into the thorax. The number to the right in each box (expressed as a percentage) is the standard error of the estimate with respect to the distance between the marker and mitral valve.

The data in Table II show the height weight body surface area sternum length thorax width and depth at mid sternum level for the 10 patients listed in the same order as Table I.

For each patient position the vertical distance of the base of the mitral valve below the marker (1) was calculated and a linear regression analysis was performed for height weight body surface area sternum length thorax width and depth. For each of the 6 positions the mean distance below the marker (average for all patients) together with the standard error of the estimate (SEL)

$$SEL = \frac{1}{N} \sum_{i=1}^N [(1 - \hat{r}_i)^2]^{\frac{1}{2}} \quad (1)$$

is shown in Table III. Next to each entry is the improved standard error of the estimate resulting from the linear regression analysis.

Discussion

For the patient in supine position a comparison between our results and those of previous studies can be made. For example Lyons and colleagues¹ placed their patients in a supine position on a rigid surface and chose the zero level as 10 cm above that surface. Richards and associates² and Cournard and co workers³ placed the level as 20 cm and 66 cm respectively below the Angle of Louis (angle between the manubrium and the anterior surface of the sternum). Weil and associates⁴ and Burri

Table 1 Location of the base of the mitral valve*

| Patient no Coordinates† | On right side 90 (cm) | On right back 45 (cm) | Flat on back (cm) | On left back 45 (cm) | On left side 90 (cm) | Tilted 30° head up (cm) |
|----------------------------|-----------------------------|-----------------------------|-------------------------|----------------------------|----------------------------|-------------------------------|
| 1 x | 65 | 73 | 36 | 48 | 43 | 40 |
| y | 49 84° | 54 08° | 33 60° | 27 24° | 41 62° | 18 34° |
| -z | 53 | 45 | 77 | 70 | 70 | 76 |
| 2 x | 65 | 67 | 43 | 61 | 57 | 73 |
| y | -11 52° | 15 50° | 23 75° | 27 18° | 27 12° | 14 78° |
| -z | 69 | 66 | 82 | 61 | 57 | 73 |
| 3 x | 59 | 69 | 62 | 71 | 64 | 64 |
| y | 07 91° | 43 11° | 38 44° | 37 41° | 30 67° | 42 55° |
| -z | 83 | 65 | 56 | 71 | 67 | 59 |
| 4 x | 64 | 48 | 45 | 57 | 43 | 17 |
| y | 24 29° | 41 10° | 43 49° | 18 28° | 48 60° | 34 15° |
| -z | 70 | 82 | 79 | 67 | 76 | 88 |
| 5 x | 47 | 52 | 37 | 46 | 34 | 43 |
| y | 27 43° | 65 61° | 14 59° | -03 16° | -08 58° | 20 39° |
| -z | 74 | 43 | 76 | 66 | 77 | 67 |
| 6 x | 39 | 34 | 34 | 37 | 36 | 48 |
| y | 10 31° | 20 61° | 16 76° | 60 27° | 23 59° | 42 37° |
| -z | 85 | 93 | 90 | 88 | 90 | 80 |
| 7 x | 48 | 58 | 28 | 42 | 32 | 44 |
| y | 10 01° | 36 24° | 34 73° | 36 34° | 37 13° | 34 67° |
| -z | 80 | 70 | 88 | 80 | 74 | 91 |
| 8 x | 41 | 41 | 36 | 55 | 42 | 43 |
| y | 50 07° | 38 06° | 46 47° | 52 64° | 44 80° | 37 56° |
| -z | 108 | 92 | 94 | 85 | 100 | 95 |
| 9 x | 50 | 37 | 38 | 39 | 30 | 20 |
| y | 06 59° | 16 30° | 22 11° | 35 83° | 32 84° | 27 41° |
| -z | 59 | 63 | 61 | 66 | 64 | 70 |
| 10 x | 67 | 57 | 46 | 46 | 35 | 58 |
| y | 38 20° | 19 55° | 26 07° | 20 26° | 28 23° | 19 10° |
| -z | 62 | 72 | 75 | 74 | 76 | 73 |

*Measured with respect to the marker placed at mid sternum

†The +x axis points toward the feet, the +y axis toward the left arm, the -z axis into the thorax

‡Rej = 1.96 standard error of the estimate (SEE) with respect to total distance from the marker to the base of the mitral valve

Clinical and analytic methods

While the analysis which forms the basis for the clinical methods is somewhat involved, the basic concepts are not difficult. By way of outline these basic considerations are summarized below. A more detailed description is in the Appendix.

In order to most simply determine the vertical distance of the mitral valve below the marker (necessary to define a reference point and located at mid sternum) a single calibrated x-ray in a horizontal projection is all that is required. A single photo necessarily "sees" only a two dimensional projection and therefore one coordinate is lost. As all three coordinates are desired, the missing coordinate is recovered by repositioning the camera.

In practice the situation is not so ideal. Due to the limited viewing field of the x-ray camera, quite often a simple horizontal projection does not include the image of both the valve and the marker. This requires viewing angles which do not allow simple inspection of the photos to reveal physiological distances with respect to a convenient patient coordinate system. In addition, the distance between the x-ray source and patient and between the patient and camera varies as the camera is repositioned, thereby changing the scale factor of every photo.

Hence an analysis of the geometry is required. That is (1) determine the camera position with respect to the patient coordinate system, (2) determine from actual

possibly to an extent not related to the quantities used here for the regression analysis

Height results in the greatest reduction in SEE in RS — 45 degrees frontal and tilt but in the remaining three positions it is last or next to last. Likewise thorax width results in the greatest improvement in the RS — 90 degrees LS — 45 degrees LS — 90 degrees positions but is last or next to last in the remaining three. Weight in every case except one (LS — 90 degrees) is third place with respect to reduction in SEE. Surface area and thorax depth vary between second and fourth place. Sternum length is last or next to last in every position except one and clearly emerges as the one parameter least likely to correlate with mitral valve location.

It is the opinion of the author that this degree of improvement in the SEE does not justify the use of a linear regression equation to estimate the base of the mitral valve for adults in the range of height weight etc covered in this study. However the use of the mean values indicated in Table III is justified as a guideline for the positioning of the zero level in adults in the positions indicated.

While no data are presented for the patient sitting upright some estimate can be made. For the patient supine (flat on back) the x coordinate (distance from marker to BMV in the direction of the feet) averages 40 cm. Tilted at 30 degrees this distance increases to 45 cm. As it appears that the heart is well tethered within the thoracic cavity we do not expect this distance to increase appreciably as the patient sits fully upright therefore we estimate in the fully upright position the BMV is 5 cm below the central sternum marker.

As stated above the main purpose of this investigation was to find more exactly the zero level for left heart pressures. A stimulus for this work was the finding in our intensive care units that continuous monitoring of left atrial pressure (LAP) after open heart surgery showed remarkable changes in this parameter when patients were turned or tilted. This happened even after readjusting the transducers according to the rules of Richards and co

workers.² Our studies show that this method is indeed misleading especially when patients were turned on their right side (Table III). Whether the implementation of our guide lines will reduce the described changes in LAP still has to be proved. In any event their use will yield more insight into the hemodynamic effects of position. Furthermore the use of Swan Ganz balloon catheters has shown the value of pulmonary capillary wedge pressure in establishing the prognosis of patients with myocardial infarctions. If this parameter (or end-diastolic pulmonary artery pressure or left atrial pressure directly) is monitored continuously for trend analysis it is necessary to know where to place the transducers in every position of the patient who will be moving much more frequently than after an operation.

Summary

The location of the base of the mitral valve was determined in 10 adult patients using a radiologic technique. The mean vertical distance below a marker placed at mid sternum was for the patient

| | |
|----------------------------|----------|
| on right side | -2.1 cm |
| | (- = up) |
| right side—back 45 degrees | 2.4 cm |
| flat on back | 7.8 cm |
| left side—back 45 degrees | 7.3 cm |
| left side | 3.0 cm |
| tilted head up 30 degrees | 8.9 cm |

The standard error of the estimate is less than 2 cm in all positions. For the patient sitting upright we estimate the distance to the BMV to be 5.0 cm below the mid sternum marker.

While some improvement in the patient to patient error could be made through linear regression based on height weight etc the degree of improvement—42 per cent or less reduction in standard error of the estimate—would not justify its use in clinical practice. The above guidelines however do provide a useful guide for setting the zero level at the base of the mitral valve.

The author thanks Mr A. den Boer for his assistance in obtaining the data and Miss I. de Wit for assistance in the analysis of the data. The authors also gratefully acknowledge the voluntary multiple retyping of this manuscript by Mrs Joan Brower.

Table III Vertical distance from marker to base of mitral valve and results of linear regression analysis

| | Patient position | | | | | |
|---|------------------|---------------|--------------|--------------|--------------|------------|
| | On right side | On right back | Flat on back | On left back | On left side | Tilted 30° |
| Mean vertical distance to mitral valve (cm) | -2.1 | 2.4 | 7.8 | 7.3 | 3.0 | 8.9 |
| Uncorrelated SEE | 1.9 | 2.0 | 1.1 | 1.7 | 1.5 | 1.0 |
| SEE correlated with height | 1.9 | 1.6 | .8 | 1.3 | 1.5 | .6 |
| % reduction SEE | 5% | 18% | 31% | 19% | 7% | 33% |
| SFE correlated with weight | 1.9 | 1.7 | .9 | 1.2 | 1.4 | .8 |
| % reduction SFE | 1.2% | 12% | 21% | 31% | 2.4% | 21% |
| SFE correlated with surface area | 1.9 | 1.7 | .85 | 1.2 | 1.5 | .7 |
| % reduction SFE | 1.2% | 15% | 29% | 30% | 1.2% | 30% |
| SFE correlated with sternum length | 1.9 | 1.9 | 1.1 | 1.2 | 1.5 | .9 |
| % reduction SFE | 0% | 5.5% | 3% | 28% | 1.1% | 8.9% |
| SFE correlated with thorax width | 1.8 | 1.8 | 1.0 | 1.0 | 1.3 | .9 |
| % reduction SFE | 4.7% | 8.9% | 7.4% | 42% | 9.7% | 2.7% |
| SFE correlated with thorax depth | 1.9 | 1.7 | 1.0 | 1.0 | 1.5 | .9 |
| % reduction SFE | 2.1% | 16% | 8.6% | 36% | 1.2% | 4.1% |

and Allgower⁵ allow for patient to patient variation by using respectively 1/2 and 3/5 of the thoracic diameter measured from the dorsal surface to the Angle of Louis. Debrunner and Buhler⁶ compared the resulting venous pressure for these various techniques for zero level determination in 26 patients undergoing right heart catheterization. Except for the method used by Lyons and colleagues,¹ all agreed acceptably well.

Our data in Table III show that the base of the mitral valve (BMV) is 7.8 cm (SEE 1.1 cm) dorsal to the mid sternal marker for the patient in the supine position. When the measurement is made with respect to the dorsal surface the distance is 13.6 cm, showing that the method of Lyons and co-workers¹ results in an overestimate of left heart pressures by about 4 cm H₂O which is consistent with the findings of Debrunner and Buhler.⁶ Furthermore, correlation with respect to thoracic depth results in only a small reduction (8.6 per cent) in standard error (Table III). Therefore those rules which place the zero level as some fraction of thorax depth will be no more accurate in adults than some fixed distance measurement. However, if this is done, our data indicate that the estimate of the BMV is 0.64 times thorax depth

measured from the dorsal surface. This is in accordance with the work of Burni and Allgower.⁵

For patients in positions other than supine there appears to be no literature with respect to the position of the BMV and the data shown in Table III represent the first report on this subject.

It was originally thought that a linear regression analysis might substantially reduce the standard error of the estimate. Unfortunately, while linear regression analysis provides some improvement in the standard error of the estimate in every case, the improvement is not especially remarkable. The improvement is greatest in the RS - 45 degrees * frontal LS - 45 degrees and tilted positions (about 18 per cent, 31 per cent, 42 per cent, and 35 per cent reduction in SEE) while the SEE in the RS - 90 degrees and LS - 90 degrees positions show virtually no reduction at a (4.7 per cent and 9.7 per cent). In the RS - 90 degrees and LS - 90 degree positions the thoracic cavity is somewhat more deformed than in the other positions—

*These abbreviations represent the following: patient positions: RS-90 patient fully on right side; RS-45; partially on right side and right back, forming an angle of 45 degrees; F flat on back; LS-45 partially on left side and back, forming an angle of 45 degrees; LS-90 fully on left side.

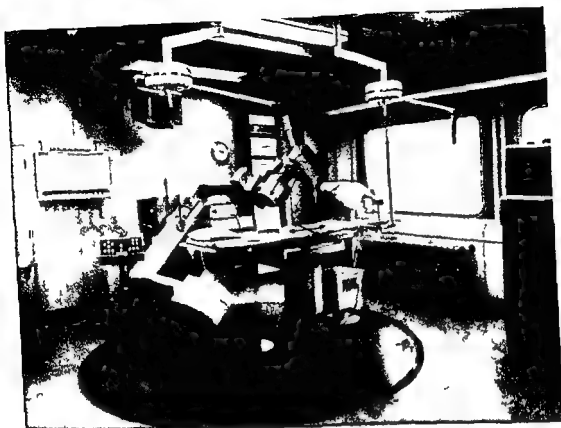


Fig 1 The apparatus used to measure position of the base of the mitral valve is shown. The arrows indicate the degrees of freedom available and the C arm angle ϕ and table angle ψ are defined. The control panel shown on the left side controls all motion except the limited horizontal motion of the table top. The x ray photo image is recorded on film and is also stored on a video disk and displayed on the monitor screen.

III Camera position A marker placed midway on the sternum (shown in Fig 2) forms the center of the coordinate system shown in Fig 3. The +x axis points toward the feet +y to the left arm and +z out of the thorax. This convention is maintained regardless of the actual patient position with respect to the x ray table and C arm and is referred to here as the patient coordinate system. The camera position is expressed by the unit vector \hat{O} with respect to the patient coordinate system as shown in Fig 3.

As shown in Fig 1 the camera position is originally recorded as the C arm angle ϕ and table angle ψ . This information together with the patient position (frontal left side 45 degrees etc) is sufficient to define the camera position with respect to the patient coordinate system. Mathematically this is simply a transformation from one coordinate system to another.

For a simple rotation about the patient's x axis (longitudinal axis) the camera position is given by equation (3)

$$\hat{O} = -\sin \phi \sin \psi \hat{i} + (\sin \phi \cos \psi \cos \gamma + \cos \phi \sin \gamma) \hat{j} + (\cos \phi \cos \gamma - \sin \phi \cos \psi \sin \gamma) \hat{k} \quad (3)$$

γ represents the patient rotation angle (0 frontal +90 lying on right side -90 on the left side etc). If the patient is tilted (rotation about his y axis) the camera position is given by equation (4)

$$\hat{O} = -(\sin \phi \sin \psi \cos \gamma + \cos \phi \sin \gamma) \hat{i} + \sin \phi \cos \psi \hat{j} + (\cos \phi \cos \gamma - \sin \phi \sin \psi \sin \gamma) \hat{k} \quad (4)$$

γ represents the tilt angle (0 for flat on back +45 if tilted head up at 45)

IV Valve location Equations (3) and (4) presents the camera location (a unit vector) with respect to the patient coordinate

REFERENCES

- 1 Lyons R L, Kennedy J A and Burwell C J. The measurement of venous pressure by the direct method. *Am Heart J* 16: 675 1938
- 2 Richards D W, Cournard A, Darling R C, Gillespie W H and Baldwin I D. Pressure of blood in the right auricle in animals and in man. Under normal conditions and in right heart failure. *Am J Physiol* 136: 115 1942
- 3 Cournard A, Riley R I, Bradley S E, et al. Studies of the circulation in clinical shock. *Surgery* 13: 964 1943
- 4 Weil M H, Shubin H and Rosoff L. Fluid repletion in circulatory shock. Central venous pressure and other practical guides. *JAMA* 192: 668 1965
- 5 Burri C and Allgower M. Klinische Erfahrungen mit der Messung des Zentralen Venen Drucks. *Schweiz Med Wochenschr* 97: 1414 1967
- 6 Debrunner E and Buhler E. Normal venous pressure, significance of reference point and normal range. *Br Med J* 3: 148 1969

Appendix

I Apparatus The system described here consisting of a floating table, C arm, and rotating floor was designed in house and is believed to be unique. Fig 1 shows the table and C arm, image intensifier, control panel and TV monitor. The arrows superimposed on the photograph illustrate the degrees of freedom available. The camera position is defined by the C arm angle ϕ and table angle ϕ_1 as shown in Fig 1.

The image intensifier is a 6 inch Philips unit and a 70 mm Philips camera is used for recording the image. With the exception of limited horizontal motion (30 cm) of the table top, all motion is powered by electric motors and is controlled at the control panel. As the photo is taken the image is simultaneously stored on a video disk. This picture is instantaneously available to verify that the exposure is correct and that the image is suitable for analysis.

There are a number of safety features incorporated whereby power is turned off when parts of the C arm come in contact with the table or patient.

II Patient procedure and scale factor determination The patient is placed on the table and raised to a height permitting positioning of the C arm. A lead marker in the shape of an 'X' is placed midway on the sternum. With the C arm in the 90 degrees position (a horizontal projection) the position of the artificial mitral valve is determined and an ink mark is placed on the side of the chest at the level of the valve. The C arm is returned to the 0 degree position (vertical) and the patient is moved horizontally out of the field of the x ray image. A square centimeter grid is then placed at the level of the mitral valve and an x ray

is taken. The patient is moved back into the x ray image and a photo is obtained of the ball valve. When the data was analyzed this information was used for measuring the ball diameter and therefore the scale factor of each photo. That is, as described above, the scale factor for each subsequent photo is not the same; the ball diameter is measured in each photo and is used to establish the scale factor.

Fig 2 illustrates the measurement procedure employed in the analysis of a single photo. The photograph distances R_1 and R_2 were measured as well as the photograph ball diameter D (averaged in two directions). Since the actual diameter of the ball is known from the grid measurement (call this diameter D_a), the physiological distance is

$$\text{physiological } R_1 = R_1 D_a / D \quad (1)$$

and

$$\text{physiological } R_2 = R_2 D_a / D \quad (2)$$

These two components constitute the vector \vec{M} described below.

The patient was placed in one of six positions: on right side (90 degrees rotation about longitudinal axis), right back (45 degrees rotation), flat on back, left back (45 degrees rotation), left side (90 degrees rotation), and tilted head up (30 degrees measured at sternum). For each position three photos were taken at different camera positions. In practice these three camera locations were never allowed to be coplanar; however, this is not an essential theoretical restriction but represents a practical consideration of weighting more equally all three dimensions.

abc matrix

$$\begin{pmatrix} 1 - a_1^2 & -a_1b_1 & -a_1c_1 \\ -a_1b_1 & 1 - b_1^2 & -b_1c_1 \\ -a_1c_1 & -b_1c_1 & 1 - c_1^2 \\ 1 - a_2^2 & -a_2b_2 & -a_2c_2 \\ -a_2b_2 & 1 - b_2^2 & -b_2c_2 \\ -a_2c_2 & -b_2c_2 & 1 - c_2^2 \end{pmatrix}$$

M matrix

$$\begin{pmatrix} x \\ y \\ z \end{pmatrix} = \begin{pmatrix} M_{x1} \\ M_{y1} \\ M_{z1} \\ M_{x2} \\ M_{y2} \\ M_{z2} \end{pmatrix}$$

(9)

It is desired to minimize the error ϵ^2 of equation (10)

$$\epsilon^2 = \sum_{n=1}^N (m_n - \alpha x_n - \beta y_n - \gamma z_n)^2 \quad (10)$$

where m_n is defined as the n^{th} element in M matrix ($m_1 = M_{x1}$, $m_2 = M_{y1}$, $m_3 = M_{z1}$, $m_4 = M_{x2}$, etc.) and α is the n^{th} element of the second row and γ the n^{th} element of the third row of the abc matrix.

Minimizing equation (10) with respect to x , y and z results in equation (11) after some rearrangement

$$x \sum_{n=1}^N \alpha^2 + y \sum_{n=1}^N \alpha \beta + z \sum_{n=1}^N \alpha \gamma = \sum_{n=1}^N \alpha m_n$$

$$x \sum_{n=1}^N \alpha \beta + y \sum_{n=1}^N \beta^2 + z \sum_{n=1}^N \beta \gamma = \sum_{n=1}^N \beta m_n$$

$$x \sum_{n=1}^N \alpha \gamma + y \sum_{n=1}^N \beta \gamma + z \sum_{n=1}^N \gamma^2 = \sum_{n=1}^N \gamma m_n \quad (11)$$

This equation contains 3 unknowns (x , y , z) which can now be solved by standard techniques. This results in a solution to equation (8) in terms of the solution generating the minimum mean square error.

In addition to the results x , y , z being printed out, the rms error ($\epsilon^2 / (N - 1) \cdot 16$) is also calculated. This has proved invaluable in detecting data handling blunders (e.g. misplaced decimal point, misreading 2 for 3, 4 for 9, etc.) and is a measure of the confidence one may place on the result.

V. Summary of methodology. A computer program was written according to the outline given below.

A. READ DATA

1. patient position (γ) x or y axis rotation
2. number of photos analyzed for a given patient position
3. arm and table angles
4. photo distance measurements

B. PRELIMINARY CALCULATIONS

1. camera location with respect to patient coordinates
2. M coefficient matrix equation (9)
3. abc coefficient matrix equation (9)
4. coefficients for 3 x 3 matrix equation (11)

C. MATRIX SOLUTION

1. solve equation (11)

D. OUTPUT

1. x , y , z coordinates
2. ($\epsilon^2 / (N - 1) \cdot 16$)
3. % error



Fig. 2 An example of a photo of the artificial mitral valve together with distance measurements made from the photo. The large X is the mid-sternum marker.

system. More briefly this can be written as equation (5) where the coefficients a , b , and c are defined by equations (3) and (4). This camera location is illustrated in Fig. 3.

$$\hat{O} = a\hat{i} + b\hat{j} + c\hat{k} \quad (5)$$

The mitral valve position is defined by another vector \hat{R} .

$$\hat{R} = x\hat{i} + y\hat{j} + z\hat{k} \quad (6)$$

as described in Fig. 3. x , y , and z are the actual physiological distances along the unit vectors defining valve location.

Viewing from the camera location \hat{O} , a two-dimensional vector \hat{U} is presented which is the measured heart position as seen from a single x-ray photo. \hat{U} is given by the vector equation

$$\hat{U} = \hat{R} - (\hat{R} \cdot \hat{O}) \hat{O} \quad (7)$$

In order to obtain the third dimension (which is collinear with \hat{O}) at least one other photo from a different observer position must be obtained. These different camera positions will be denoted by subscripts 1, 2, etc. (in practice 3 different observer angles were used to allow for the

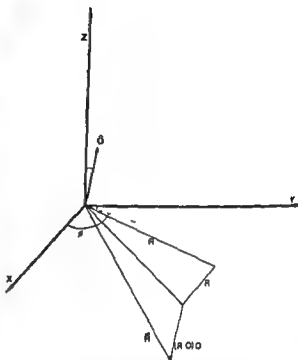


Fig. 3 Patient coordinate system used in the analysis of the data. The origin is at the mid-sternum marker. The $+x$ axis points towards the feet, $+y$ axis towards the left arm, and $+z$ axis out of the thorax.

contingency of some photos being spoiled during exposure or processing). The equations which must be solved to determine heart position is summarized by equations (8).

$$\begin{aligned} \hat{U}_1 &= \hat{R} - (\hat{R} \cdot \hat{O}_1) \hat{O}_1 \\ \hat{U}_2 &= \hat{R} - (\hat{R} \cdot \hat{O}_2) \hat{O}_2 \end{aligned}$$

$$\hat{U}_n = \hat{R} - (\hat{R} \cdot \hat{O}_n) \hat{O}_n \quad n \geq 2 \quad (8)$$

For $n \geq 3$ these equations appear to over-specify the problem and additional photos (more than 2) would not appear to be necessary nor desirable. In fact given actual patient data no exact solution to equations (8) exists due to various combinations of measurement and round off errors and actual motion of the heart. Given this fact a solution is sought on the basis of minimum least squares error. This criterion can of course be applied to two photos but greater confidence in the result is obtained with a larger number.

Expressing equations (8) in matrix notation results in equation (9).

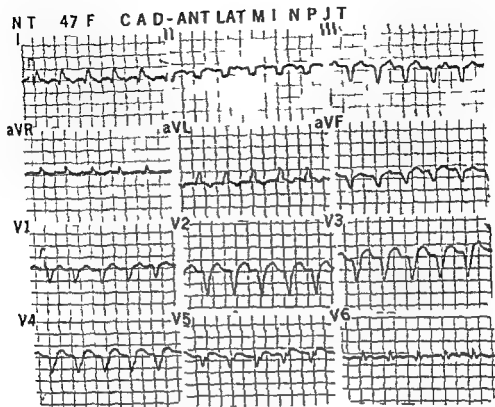


Fig. 1 Twelve-lead electrocardiogram recorded in a 47-year-old woman with sinus tachycardia, A-V dissociation (rate = 104 per minute) and an acute anterior wall myocardial infarction. The QRS complexes are prolonged at 0.13 second; their appearance was identical during intact A-V conduction.

and 18 women whose ages ranged from 41 to 87 years with a mean age of 64 years. When subdivided according to location of infarction as determined by usual electrocardiographic criteria, there were 24 cases of inferior wall infarction (30 per cent), 16 cases of anterior wall infarction (33.3 per cent), and eight cases of infarction at other sites (posterior isolated lateral 16.5 per cent). Congestive heart failure was classified as mild when there were bibasilar rales without dyspnea, moderate when rales were heard over the entire lower half of the anterior and posterior thorax and severe in the presence of pulmonary edema. Patients with significant diminution of blood pressure in the absence of any acute postural change or administration of narcotics were classified as having either hypotension (a decline of brachial blood pressure greater than 70 mm Hg in systolic and diastolic levels) or shock when the drop in blood pressure was accompanied by

cold diaphoretic extremities, a weak pulse and a decline in urinary output. In those cases where the electrocardiographic differential diagnosis was accelerated ventricular rhythm vs. VPJT with aberrant ventricular conduction, the latter diagnosis was accepted only if there were identifiable sinus capture beats with QRS complexes identical with those seen during the tachyarrhythmia. Only four cases fell into this latter category. Significance of the difference between mean values was determined using Student's *t* test.

Results

Type of non paroxysmal A-V junctional tachycardia. The most prevalent form of VPJT was not associated with identifiable independent atrial activity (28 cases, 58 per cent). The balance of the types of VPJT encountered were as follows: complete A-V dissociation (six cases, 12.5 per cent, Fig. 1); incomplete A-V dissociation

Non-paroxysmal A-V junctional tachycardia associated with acute myocardial infarction

Jaco Iishenfeld, M D
Kenneth B. Desser, M D
Alberto Benchimol, M D
Phoenix, Ariz

Non paroxysmal A V junctional tachycardia (NPJT) results from an acceleration of impulse formation at the region of the A V junction¹. This tachyarrhythmia is usually associated with digitalis toxicity, acute rheumatic fever or inferior wall infarction¹ and generally produces a ventricular rate of 60 to 150 beats per minute.² Despite existing evidence that this dysrhythmia does not per se indicate a poor prognosis in the setting of acute myocardial infarction,^{1,2} a recent study has demonstrated a high mortality rate in subjects with acute infarction and NPJT.³ Since our impression based on clinical experience did not correspond with that of the recent study, we reviewed electrocardiographic findings and clinical outcome in a group of patients with NPJT which occurred in the setting of acute myocardial infarction.

Materials and method

The records of all subjects who were admitted to the Good Samaritan Hospital with the provisional diagnosis of acute myocardial infarction during the years 1970 and 1971 were reviewed. These years

were selected because of the profusion of rhythm strips added to permanent clinical records during admission to the Coronary Care Unit. Patients were accepted as having acute myocardial infarctions when their electrocardiograms demonstrated classical significant Q waves with evolving ST segment and T wave changes compatible with that diagnosis associated with a history of at least one hour of retrosternal chest pain. Furthermore, all subjects had serial elevations of serum glutamic oxaloacetic transaminase (SGOT) to at least twice control levels. The normal range for SGOT at this institution is 0 to 50 IU. Based on the above criteria, 49½ patients had acute myocardial infarction. Of these, 45 subjects (91 per cent) had NPJT. The records of these patients were reviewed specifically for the purpose of determining survival at peak SGOT level, ventricular rate during NPJT, day of occurrence of the tachycardia and symptoms produced by it. Subjects with evidence of digitalis toxicity and those who were receiving atropine or adrenergic stimulating drugs prior to the onset of the tachyarrhythmia were excluded from the study. There were 30 men

From the Institute for Cardiovascular Diseases, Good Samaritan Hospital, 1033 East McDowell Rd., Phoenix, Arizona 85006.

Supported in part by the Nichols Memorial Fund.

Received for publication Jan. 22, 1973.

Reprint requests to Alberto Benchimol, M.D., Good Samaritan Hospital, P.O. Box 2989, Phoenix, Arizona 85062.

December 1973

course of acute myocardial infarction was significantly higher in those patients who died (survivors 152 ± 114 non survivors 458 ± 319 $P < 0.01$)

Onset of NPJT and relationship to symptoms In the vast majority of cases (84 per cent) NPJT appeared during the first three days of hospitalization. The time of onset in the non survivors paralleled that of the entire study group with 80 per cent of cases manifesting this arrhythmia during the initial 72 hours of observation. No episode of acute pulmonary edema hypotension or shock was produced or worsened by the occurrence of NPJT. Mild to moderate congestive heart failure occurred as a direct consequence of NPJT in six subjects. The respective locations of infarction in this latter group were three cases—anterior three cases—inferior. The average ventricular rates in these six patients tended to be higher (mean—132 per minute) than rates in those subjects in whom symptoms were not evoked by NPJT (mean—90 per minute).

Discussion

The underlying basis of accelerated A V junctional discharge associated with acute myocardial infarction is not well established.³ Local effects of increased catecholamine and potassium levels⁴ have been advanced as possible reasons for increased A V junctional automaticity in the setting of acute infarction yet there is no substantial proof for such operative mechanisms. The incidence of NPJT in acute infarction and the relative distribution of anatomic sites described in this study is comparable with the data of Konecke and Knoebel² who found 10 per cent of 203 subjects with coexisting acute infarction and NPJT.

Furthermore NPJT occurred almost twice as frequently with inferior wall as compared with anterior wall infarction in their study. Examination of the respective mortality rates described by Konecke and Knoebel and those set forth herein differ in that we found only a single subject with inferior wall infarction and NPJT who died. In contrast the other authors described a mortality rate of 43 per cent under similar circumstances. In general a poor prognosis can be assigned to those

subjects with anterior wall infarction and NPJT since the mortality rate in patients with this combination of findings was previously found to be almost 100 per cent² and was 62 per cent in this study.

In our experience NPJT per se rarely produces a change in the clinical status of subjects with acute myocardial infarction. The major determinants for ultimate clinical outcome appear to be the ventricular rate during NPJT and peak ST OT level. It is possible that the more rapid ventricular rates associated with NPJT increased the extent of myocardial damage⁵ and consequently the mortality rate due to shock power failure asystole and secondary ventricular fibrillation. Alternatively the more rapid A V junctional discharge rates may have been a reflection of greater underlying myocardial necrosis. The presence of significantly higher peak ST OT levels in those patients with NPJT who succumbed is in accord with recent evidence which suggests that in general the extent of myocardial infarction can be grossly reflected by the magnitude of serum enzyme rise.⁶ Over all mortality rates during acute infarction appear to have a highly significant linear correlation with the ST OT level.⁷ If these observations apply to the data presented in this study then it could be surmised that the extent of myocardial necrosis in patients with anterior wall infarction and coexisting NPJT was of a greater magnitude than that in subjects with inferior wall infarction and identical dysrhythmias. Detailed clinicopathologic study appears necessary in order to shed light on this possibility.

In conclusion NPJT does not necessarily indicate a poor prognosis in the setting of acute myocardial infarction. Our data do suggest however that the mortality rate in subjects with coexisting NPJT and anterior wall infarction is high.

Summary

The hospital records of 48 subjects with acute myocardial infarction complicated by non paroxysmal A V junctional tachycardia (NPJT) were reviewed. Fifteen of 48 subjects (31 per cent) so affected died. NPJT was most commonly associated with inferior wall infarction (24/48 50 per cent)



Fig. 2 Variations of P wave configuration noted in a 54 year old man with acute inferior wall myocardial infarction prior to established NPJT. Lead II rhythm strip demonstrates gradual change in P wave polarity from upright to total inversion.

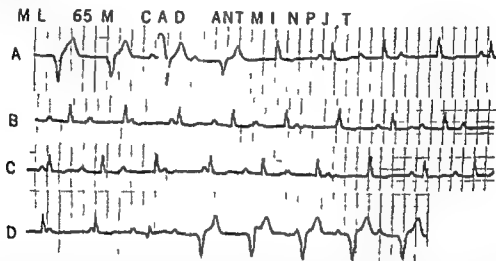


Fig. 3 I through D Lead II rhythm strip from a 65 year old man with an acute anterior wall myocardial infarction sinus rhythm complete AV block NPJT (rate = 10 per minute) and intermittent accelerated ventricular rhythm. Widened negative QRS complexes occur at the beginning of record 1 and end of record D; their intrinsic rate is slightly more rapid than that of the NPJT. Fusion beat (fourth QRS complex in record 1; third QRS complex in record D) can be seen. The wide QRS complexes presumably arise from an accelerated ventricular (or fascicular) subsidiary pacemaker which usurps control of the ventricles.

(seven cases 14.3 per cent) wandering atrial pacemaker (five cases 10.3 per cent Fig. 2) and retrograde AV junctional Wenckebach phenomenon (two cases 4 per cent). In a single patient apparent episodes of accelerated ventricular rhythm occurred in the setting of complete AV block and NPJT (Fig. 3). There was no relationship between the site of infarction and the type of NPJT. Furthermore, the variations of NPJT did not correlate with survival.

Mortality and relation to site of infarction

Of 48 patients 15 died (31 per cent). Ten of 16 subjects (62.5 per cent) with anterior wall myocardial infarction died during the course of NPJT. On the other hand 23 of 24 patients (96 per cent) with inferior wall infarction survived. Four of eight (50 per cent) subjects with infarctions at other sites died. There were no significant differ-

ences between the mean ages or number of previous infarctions in the survivor and non-survivor groups.

Ventricular rate. Mean (± 1 standard deviation) heart rates during NPJT in patients with inferior and anterior wall myocardial infarctions were 87.4 ± 25.1 and 113.4 ± 37.3 beats per minute respectively. The difference between these values was statistically significant ($P < 0.01$). Six of ten patients (60 per cent) with anterior wall myocardial infarction who died had ventricular rates greater than 100 per minute during NPJT. The only subject with inferior wall infarction who did not survive had a heart rate of 110 per minute. Nine of all 15 subjects (60 per cent) who died manifested rates equal to or greater than 100 per minute during NPJT.

Peak SGOT levels. The mean value for peak SGOT level obtained during the

Clinical diagnosis of persistent left superior vena cava by observation of jugular pulses

Simón Horuit^a M D

José Esquivel A M D

Faust Altie M D

Eulo Lupi H M D

Jorge Espino Vela M D

Mexico City Mexico

The persistence of the left superior vena cava (LSVC) is a congenital anomaly of the coronary sinus usually draining through this vessel. It has no hemodynamic importance.^{1,4} In 3 to 4 per cent of the cases it is an association of other congenital cardiovascular malformations.⁴ The persistence of LSVC may be important for the surgeon because it requires special handling in extracorporeal circulation.¹

Generally the diagnosis is made by intra cardiac catheterization or by surgery. The possibility of a clinical diagnosis on the basis of the radiological appearance^{2,4,6} and by the deviation of the mean electrical axis of the P wave toward the left has been reported.^{4,7,8}

In 1967 Colman¹⁰ described a case in which the pulse of the left jugular vein was more prominent than that of the right one and this was considered a sign of persistence of LSVC. This finding has not been mentioned again in the literature. For this reason we investigated it in patients in whom the malformation was proved in an attempt to elicit its usefulness and incidence.

Methods

The study included six patients in whom an LSVC had been demonstrated by intra cardiac catheterization or at surgery and a seventh case with mirror image dextrocardia with situs inversus and persistent right superior vena cava. A difference in the amplitude of the external jugular pulses in both sides of the neck was looked for. The jugular phlebogram was recorded subcutaneously using a four channel jet inscription polygraph (Mingograph 34 Elema Schonander). Simultaneous recording of an ECG lead and a phonocardiogram with microphone EMT 25 and logarithmic filter were obtained. The pressure phlebogram was inscribed with a Bouche Brecht Infratone receptor¹¹ with linear response of 0.02 to 200 Hz following the technique described by Hartmann.¹² The receiver was held with a pneumatic cuff with a variable pressure of 2 to 10 mm Hg. Simultaneous phlebographic bilateral recordings were obtained when possible otherwise the same pressure was applied in each one of the recordings. The same procedure was used in three patients without the anomaly. The paper

From the Instituto Nacional de Cardiología de México.

Received for publication July 22, 1973.

Reprint requests to Simón Horuit, M.D., Instituto Nacional de Cardiología, Ave. Miguel Alemán 300, México 7 D.F., México.

Although ten of 16 (63 per cent) patients with acute anterior wall infarction and NPJT died 23 of 24 patients with acute inferior wall myocardial infarction survived. Mean heart rates during NPJT were significantly greater in subjects with anterior wall infarction (113.4 ± 35.3 vs inferior wall 85.4 ± 28.1 $P < 0.01$). Peak SGOT levels were significantly higher in those patients who died (488 ± 579 vs survivors 152 ± 114 $P < 0.01$). NPJT altered the clinical status of only six subjects. It is concluded that NPJT indicates a poor prognosis in subjects with acute anterior wall infarction but is generally associated with a benign clinical course in patients with inferior infarction. These differences may be based on a greater extent of myocardial damage in the former group.

We wish to acknowledge the technical assistance of Nancy Copeland, R.N., Carole Crevier, Larry

Kuriger, Sydney Peebles, Sharon Squire, and Les Zendle.

REFERENCES

1. Pick A and Dominquez P. Non paroxysmal A-V nodal tachycardia. *Circulation* 16:1077 1957.
2. Konecke L L and Knoebel S H. Non paroxysmal junctional tachycardia complicating acute myocardial infarction. *Circulation* 45:567 1972.
3. DeSanctis R W, Block P and Hutter A M Jr. Tachyarrhythmias in myocardial infarction. *Circulation* 45:681 1972.
4. Shell W E, Henry R D and Sobel B E. The effect of increased heart rate on infarct size in the conscious dog (Abstract). *Circulation* 44(Suppl 11):61 1971.
5. Sobel B E and Shell W E. Serum enzyme determinations in the diagnosis and assessment of myocardial infarction. *Circulation* 45:471 1972.
6. Chapman B L. Relation of cardiac complication to SGOT level in acute myocardial infarction. *Br Heart J* 34:890 1972.

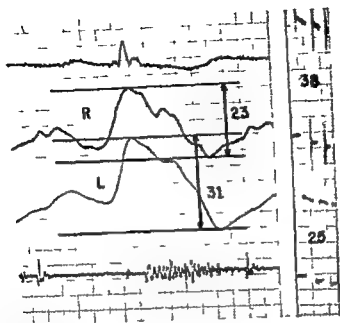


Fig 1 Jugular phlebogram of Case No. 1. Note that the height of the phlebogram has been corrected in the table according to the amplification. (ECG Lead II and phonocardiogram are on top and bottom. Numbers represent mm.)

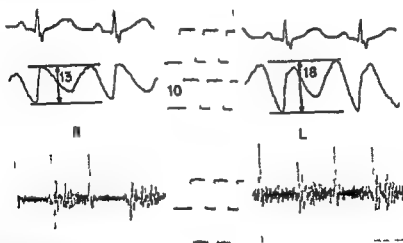


Fig 2 Jugular phlebogram of Case No. 2 (see Table I). (ECG lead and phonocardiogram are on top and bottom. Numbers represent mm.)

catheterization does not always elicit the malformation only 68 to 90 per cent of the cases are diagnosed. Its clinical diagnosis would prompt the intentional searching by catheterization or anticipate its existence thus preventing unexpected shortcomings or other problems in the introduction of a pacemaker electrode by a peripheral vein through the left arm⁸ and in modifying

techniques of cannulation for extracorporeal circulation.¹

Persistence of LSVC has been anticipated from the radiological image in 39 to 86 per cent of the cases^{4, 6, 12}—a visible shadow to the left of the cardiac contour. The left axis deviation of the P wave in 34 to 70 per cent of the cases^{4, 7, 9}—another useful indication.

The description by Colman¹⁰ of the

Table I Patient data and relationship between left and right phlebograms

| Case | Age yr | Sex | Diagnosis | Diagnosis L S V C* | Height of jugular phlebogram (mm) | | Left right ratio |
|------|-----------|-----|--|-----------------------|--------------------------------------|-------|---------------------|
| | | | | | Left | Right | |
| 1 | 17 | F | Dextroversion in situs solitus Complete transposition of the great arteries with ventricular inversion Pulmonary stenosis ventricular septal defect L S V C to C S | C | 12.4 | 6 | 2 |
| 2 | 4 | M | Atrial septal defect with pulmonary stenosis Patent ductus arteriosus (operated) I S V C to C S | S | 18 | 13 | 1.3 |
| 3 | 5 | F | Aortic incompetence and ventricular septal defect pulmonary stenosis I S V C to C S | C S | 17.5 | 12.2 | 1.43 |
| 4 | 7 | F | Ventricular septal defect Pulmonary hypertension L S V C to C S | C S | 15 | 9 | 1.66 |
| 5 | 4 | M | Transposition of the great arteries Double outlet right ventricle Pulmonary stenosis Ventricular septal defect Atrial septal defect L S V C to C S | C | 8.9 | 5.2 | 1.7 |
| 6 | 3 | F | Double outlet right ventricle Atrio ventricularis communis L S V C to C S | C | 13 | 19 | 1.46 |
| 7 | 8 | F | Mirror image dextrocardia situs inversus Patent ductus arteriosus ventricular septal defect atrial septal defect R S V C to C S | C | 20 | 32 | 0.62† |

*Abbreviations: L S V C = left superior vena cava; R S V C = right superior vena cava; C S = coronary sinus; C = catheterization; S = surgery.

†See text for explanation.

recording speed was 50 to 100 mm per second.

The distance from the peak of the α wave to the lowest point usually the x collapse was measured in each tracing. The height of the c wave was not measured since it is influenced by the pulse of the carotid artery. The height was adjusted by the amplification with the following formula:

$$\frac{\text{Height of phlebogram in mm} \times 10}{\text{Amplification in mm}}$$

A left right phlebogram ratio was calculated as follows:

$$\frac{\text{Height of left phlebogram}}{\text{Height of right phlebogram}}$$

Results

Age, sex, and associated heart disease of the cases are summarized in Table I. In six

of the seven cases the left jugular pulse had greater amplitude on visual inspection than the right one and the left right phlebogram ratio exceeded one. In the seventh case the right jugular pulse had greater amplitude demonstrated with phlebographic tracings. This patient had mirror image dextrocardia with situs inversus—i.e. there was an anomalous right superior vena cava. The three control patients showed a larger right phlebographic tracing with a left/right ratio of 0.15, 0.25, and 0.42, respectively.

The different values of the phlebographic tracings and the relation between the left and right phlebograms are described in Table I. Figs. 1, 2, and 3 are examples of this condition.

Discussion

The importance of the diagnosis of LSVC has been emphasized.^{1,2} A cardiac

- Blount G Persistent left superior vena cava
Am J Cardiol 14:77 1959
- 3 Aguilar J C, Soni J, de la Cruz M V
and Rubio V Vena cava superior izquierda
Presentacion de 42 casos Arch Inst Cardiol
Mex 30:452 1960
- 4 Horvatz S, Eparza J, Lupi H E and
Epino Vela J Diagnostico clinico de la
persistencia de la vena cava superior izquierda
Arch Inst Cardiol Mex 42:102 1973
- 5 Winter F W Persistent left superior vena
cava Angiology 5:90 1954
- 6 Zappacosta C, Rousell C, Castellino V
and Ferrane J Diagnostic radiologique de la
veine cave supérieure gauche J Fr Chir
Thorac 21:327 1967
- 7 Hancock E W Coronary sinus rhythm in
sinus venosus defect and persistent left superior
vena cava Am J Cardiol 14:608 1964
- 8 Osawa M Leftward shift of P wave axis and
its diagnostic consideration in sinus venosus
defect and persistent left superior vena cava
Jap J Thorac Surg 19:617 1966
- 9 Momma K and Linde I M Abnormal
rhythms associated with persistent left superior
vena cava Radiat Res 31:10 1969
- 10 Colman A I Diagnosis of left superior vena
cava by clinical inspection a new physical sign
Am Heart J 73:115 1967
- 11 Boucke H and Brecht K Ein neuer elek-
trischer Aufschreiber und seine einfachste an-
wendung in der ärztlichen praxis Dtsch Med
Wochenschr 88:562 1952
- 12 Hartman H The jugular venous pulse tracing
Am Heart J 49:698 1960
- 13 Fraser R S, Dworkin J, Rossall R E and
Eidem R Left superior vena cava A review
of a associated congenital heart lesions catheteri-
zation data and roentgenologic findings Am J
Med 31:711 1961
- 14 Sleight I Unilateral elevation of the internal
jugular pulse Br Heart J 24:726 1962

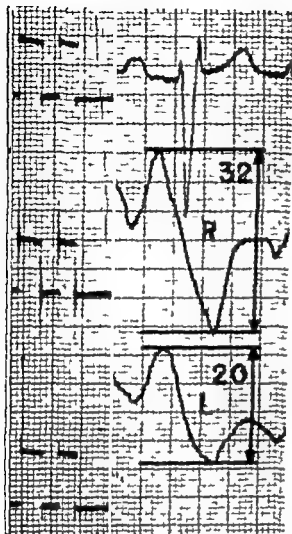


Fig 3 Jugular phlebogram of Case No. 7. The right pulse is of greater amplitude but the patient has situs inversus with persistent right superior vena cava. (See table and text. ECG Lead II is on top. Numbers represent mm.)

greater amplitude of the left jugular pulse in a proved case with LSVC was observed and demonstrated in all our cases. The prominence of the right jugular pulse in the patient with dextrocardia and situs inversus is considered positive on account of the expected inversion and this allows us to consider a new clinical evidence probably more consistent in the diagnosis of this anomaly.

The mechanisms advocated by Colman¹⁰ explain the greater amplitude of the left jugular pulse. Normally the right jugular pulse is either equal or even higher than the left one as is shown in our controls. This is due to the fact that the right venous system of the neck is in a relatively straight communication with the right atrium whereas the left internal jugular vein com-

municates with the superior vena cava via a longer and nearly horizontal channel.¹⁰ In the presence of persistent LSVC the participation of the coronary sinus during atrial systole or the greater atrial pressure on the coronary sinus may transmit the pressure wave better toward the venous system in the left side and pressure measurements on both superior vena cavae when coexisting should be performed to assess the hypothesis.

The greater amplitude of the left jugular pulse can be found in other circumstances. The obstruction or absence of the right superior vena cava can decrease the pressure in the homolateral jugular vein enough to make the pulse unequal¹⁰ phasic or expiratory obstruction of the left innominate vein by an uncoiled aortic arch can increase the pressure in the internal jugular vein¹⁴ as is seen in elderly and arteriosclerotic patients. The compression of the venous system of the left side by neoplastic processes of the superior mediastinum can also increase the venous pressure of the same side. Finally anomalous drainage of the pulmonary veins to the left innominate vein sometimes associated with persistent LSVC, could increase the amplitude of the left jugular pulse. These situations can be suspected by other signs and so the differential diagnosis is easily done.

Summary

In seven patients with persistent left superior vena cava (LSVC) diagnosed at surgery or by intracardiac catheterization the difference in amplitude between the left and the right jugular pulses was investigated. In six cases the left jugular pulse had greater amplitude and in the seventh case the opposite was true but this patient had mirror image dextrocardia with situs inversus and therefore it was considered a positive sign. The greater amplitude of the jugular pulse in the same side of the anomalous superior vena cava (usually the left) appears to be a useful sign to suspect the diagnosis of this malformation.

REFERENCES

1. Frank C G and Maloney J V. Surgical significance of congenital anomalies of the coronary sinus. *J Cardiovasc Surg* 9:420 1968.
2. Gensini C G, Caldini P, Casaccio F and

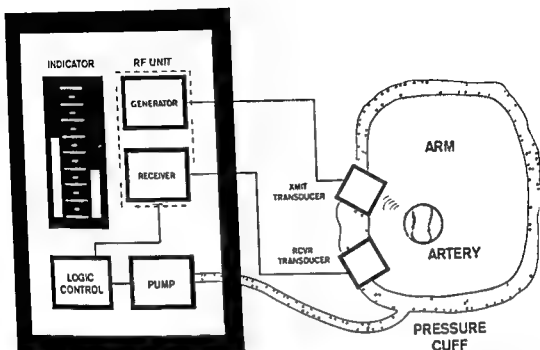


Fig 1 Principle of operation of the automatic Doppler ultrasound blood pressure system. Ultrasonic energy is generated and transmitted via the transmitter transducer into the arm toward the brachial artery. Reflected waves are sensed by the receiver transducer and receiver. An occlusive cuff is inflated to above systolic then slowly deflated. Systolic and post systolic arterial wall motion is sensed by the logic as a change (Doppler shift) in the frequency of the reflected wave. The logic determines systolic and diastolic pressures and displays them on the indicator.

ures (Fig 1). An occlusive cuff is placed on the arm in the usual manner with an ultrasonic transducer on the arm over the brachial artery. Upon operator activation of the system, the cuff is inflated automatically to a preset pressure and deflated at the rate specified by the American Heart Association.⁸ A low energy⁹ ultrasonic field is transmitted into the arm; the portion of the ultrasound that is reflected by the arterial wall shifts in frequency (Doppler shift) when the wall of the artery moves. Above systolic the vessel remains closed due to the pressure of the occluding cuff and motion signals are not received. As cuff pressure falls to the point where it is just overcome by brachial artery pressure, the artery wall snaps open. This opening wall movement often corresponding to the occurrence of the first Korotkoff sounds¹ produces a Doppler shift which is inter-

preted after several checks by logic in the instrument as systolic and causes a mercury column to lock in place at the indicated pressure. As cuff pressure declines further, the instrument continues to analyze ultrasound reflected from the arterial wall. At a point corresponding to the fourth phase of Korotkoff sounds (muffling) the violence of wall motion markedly diminishes.⁵ The instrument notes the sudden diminution in the amplitude of the Doppler shift and cuff pressure at this point is displayed as diastolic pressure. The system is totally automatic. The operator positions the cuff/transducer assembly, pushes a button and reads the results.

Since the use of Doppler ultrasound blood flow detectors in the measurement of BP has also been reported,^{8,11} the similarities and differences in the two ultrasonic approaches should be briefly noted. Both depend on motion detection by Doppler shift during cuff deflation. In a

Automatic ECG and blood pressure measurement in multitesting

Correlation of blood pressure and ECG abnormalities

H M Hochberg MD*

M L D George BS LE**

E L Schmalzbach***

C A Caceres MD****

Cranbury N J

Large health testing demonstrations allow rapid large scale testing of new concepts and instruments as well as extraction of new data inter relations. Previous demonstrations had been used to develop and validate on line computer electrocardiogram (ECG) analysis and derive ECG classifications.¹⁻⁴ This report describes the inter relations of these ECG classifications with automatically derived blood pressure. The 1108 participants in the American Dental Association Annual Health Evaluation Program (1970) had the following tests: an automated ECG by the Medical Systems Development Laboratory Program;¹⁻⁴ automatic blood pressure measured with an instrument approaching intra arterial pressure in accuracy;^{5,7} clinical chemistry (blood urea nitrogen, blood glucose, serum cholesterol and uric acid); tonometry; head, neck, and oral examina-

tion; and x ray examination of teeth, jaw and chest.

The results showed blood pressures were distributed in a manner similar to the US population and ECG classifications were similar to those reported in prior multitesting of this group. There is strong linear relation between the per cent of questionable ECG's and the systolic and diastolic pressures. Expected correlations between age and QRS axis and pressure and ECG wave form amplitudes were not found.

System description

The ECG system has been described in detail.² The blood pressure (BP) measurement system is an ultrasonic device capable of sensing arterial wall motion combined with a computer that analyzes the wall motion signals to detect systolic and diastolic and display the corresponding pres-

From the Biomedical Research Department, Roche Medical Electronics Division, Cranbury, N. J.

Received for publication Feb. 8, 1973.

Reprint requests to H. M. Hochberg, M.D., Director, Biomedical Research Department, Roche Medical Electronics Division, Cranbury, N. J. 08512.

*Director, Biomedical Research Department, Roche Medical Electronics Division.

**Engineer, Biomedical Research Department, Roche Medical Electronics Division.

***Engineering Specialist, Biomedical Research Department, Roche Medical Electronics Division.

****Consultant, Clinical Systems Associates, Inc., Washington, D. C.

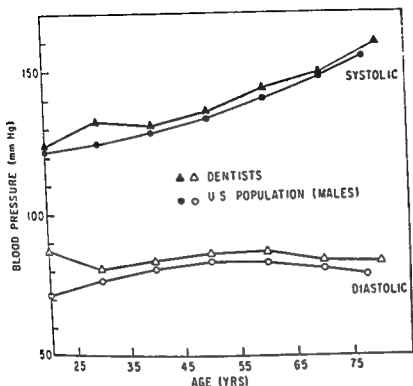


Fig 2 Blood pressures obtained at the 1970 Health Evaluation Program is compared to the U.S. population. Systolic and diastolic pressures closely paralleled the U.S. population except in the youngest group which contained only one subject. The slight difference in absolute values may be due to the difference in instruments used or observer variation. This difference is probably not clinically significant.

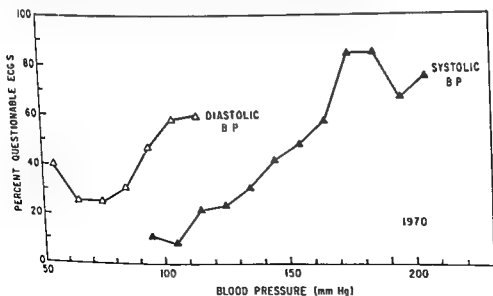


Fig 3 Relation of proportion of questionable ECGs to systolic and diastolic blood pressure. The percentage of questionable ECGs rises smoothly with both systolic and diastolic blood pressures, plateauing well in the hypertensive range. There are no definite breaks until the plateau, the proportion rising constantly with blood pressure.

Table 1 Weight blood pressure, and ECG findings in American Dental Association screening

| Age (yr) | No | Mean weight (kg) | Hypertension | | | | ECG questionable | |
|--------------|------|------------------------|--------------|------|-----------|-----|------------------|------|
| | | | Systolic | | Diastolic | | No | % |
| | | | No | % | No | % | | |
| 0-24 | 1 | 71.3 | 0 | 0 | 0 | 0 | 0 | 0 |
| 25-34 | 156 | 81 | 15 | 10 | 2 | 1 | 36 | 23.1 |
| 35-44 | 293 | 81 | 29 | 10 | 16 | 8 | 64 | 21.8 |
| 45-54 | 344 | 80 | 57 | 17 | 30 | 9 | 116 | 33.7 |
| 55-64 | 193 | 80 | 61 | 32 | 12 | 6 | 98 | 50.7 |
| 65-74 | 105 | 79 | 44 | 42 | 10 | 9 | 56 | 53.4 |
| 75-84 | 16 | 72.3 | 10 | 62 | 0 | 0 | 10 | 62.5 |
| Total (mean) | 47.5 | 1.108 | 216 | 19.5 | 40 | 6.9 | 380 | 34.4 |

blood flow detector the ultrasonic frequency is set to produce a signal ascribed to backscattering from moving blood cells.¹⁰ This instrument however uses lower ultrasonic frequency to reduce blood flow backscatter filters out the higher frequency reflections and passes the lower frequencies thus following movement of the relatively slow moving arterial wall. Clinically the major apparent difference is a reported inability of the blood flow detectors to obtain diastolic pressures in some patients during shock¹⁰ and in smaller infants.¹¹ However as previously described this instrument is an automated system and thus has advantages in screening applications where rapidity and avoidance of observer bias and fatigue are important considerations.

Method

Participants in the American Dental Association Convention (1970) were invited to the Annual Health Evaluation Program, a series of health tests. The tests included an automated electrocardiogram by the Medical System Development Laboratory Program,¹² automatic BP with the instrument described above, clinical chemistry (blood urea nitrogen, blood glucose, serum cholesterol, and uric acid), tonometry, neck, and oral examination and x-ray examination of teeth, jaws, and chest. The

ECG had been automatically analyzed at the Health Evaluation Program in previous years by the same program reported here. Results have been reported elsewhere.^{4,5}

To facilitate rapid recording of BP, two tables were prepared with the system cuff transducer and cables appropriately positioned and fastened for rapid application and removal. The operators had no previous knowledge of or experience in BP measurement and were easily trained. Computer ECG analyses were reviewed and were classified into normal and questionable based on previously reported criteria⁴ by one of the authors (C. A. Caceres).^{*} Participants' BP, age, height, weight, and ECG results were recorded. Results were subjected to computer analysis.

Timing of the blood pressure analysis portion of the exam showed that each participant was processed in an average time of 45 seconds. Each of the two instru-

Electrographic interpretations were classified as normal or questionable without knowledge of history, age, sex, or position of the participant. Tracings were considered normal if the following conditions were met: (1) the QRS axis was between -19° degrees and $+95^\circ$ degrees; (2) the heart rate was between 56 and 109 beats per minute; (3) no more than one minor abnormality was found (e.g., minimal ST depression, horizontal QRS, abnormal P wave axis, etc.); or (4) no arrhythmia was present. The others, the questionable ECGs, had findings probably of clinical significance.

Individual blood pressures are usually not considered hypertensive until a level of >160 mm Hg systolic or >100 mm Hg diastolic has been attained. These results showed 19 per cent with systolic and 6.9 per cent with diastolic hypertension. The U.S. figures based on a lower average age are 17 per cent and 5.5 per cent.¹²

The differences from the U.S. population mean for each age group are well within the bounds of observer variation^{14,15} and are thus probably not clinically significant. The small consistent difference noted may be due to the use of the more sensitive ultrasound method or to the activity of subjects attending a meeting.

As expected the blood pressure level rose with age but the body weight did not. The percentage of questionable ECG's is similar to previously reported data on a similar population of dentists.^{1,4,5} These data indicate that the population screened at the 1970 Annual Health Evaluation is representative of the U.S. male population with respect to blood pressure. We believe from unpublished data (C. A. Cicere's) that this applies to the ECG as well.

The almost linear relation of the proportion of questionable ECG's to BP was not expected. Another unexpected finding was the lack of correlations of age and BI with the ECG changes they are usually associated with. Review of the age and BI distribution shows the number of extreme cases may be too small to allow this correlation to become clear. Nonetheless the ECG does change with BI but in a non-specific way.

Summary and conclusion

An automatic ultrasound blood pressure instrument was used to measure blood pressure (BP) and a computer was used to analyze ECG's in 1108 male professionals at a voluntary Health Evaluation Program at a professional society meeting. BP's were recorded with the technique recommended by the American Heart Association in an average time of 45 seconds per participant by technicians untrained in the auscultatory method of obtaining BP. Nineteen per cent of the participants had systolic hypertension (>160 mm Hg) and 7 per

cent had diastolic hypertension (>100 mm Hg). The mean BP's and percentage of abnormalities were equivalent to the U.S. population. Computer ECG analysis showed 34.4 per cent questionable findings, consistent with previous experience. The study showed a strong positive linear relationship between the proportion of questionable ECG's and the systolic and diastolic blood pressure.

The author wishes to thank Mr H. C. Ching and Mr C. Rosencrown, Roche Medical Electronic, who performed the statistical analyses presented here.

REFERENCES

1. Haxberg, H. M., Wehrer, A. I., Abraham, S. and Cicere, C. A. Automatic ECG analysis. *J. Am. Dent. Assoc.* 73:644, 1966.
2. Hochberg, H. M., Calitry, J. B., Wehrer, A. I., Whitteman, J. and Cicere, C. A. Automatic ECG analysis in rapid mass screening. *Arch. Environ. Health* 15:390, 1967.
3. Cicere, C. A. and Haxberg, H. M. Performance of the computer and physician in the analysis of the electrocardiogram. *Am. Heart J.* 9:339, 1970.
4. Cicere, C. A. and Dreifu, L. S. Clinical electrocardiography and computers. New York, 1970. Academic Press.
5. Haxberg, H. M. and Salomon, H. Accuracy of an automated ultrasound blood pressure monitor. *Curr. Ther. Re.* 13:139, 1971.
6. Sheppard, I. C., Johnson, T. S. and Kirklin, J. W. Controlled study of brachial artery blood pressure measured by a new direct method. *J. Am. Assoc. Med. Instr.* 21:977, 1971.
7. Lopper, P. J., Fichten, R. M. and Donham, R. T. Automatic ultrasound monitoring of blood pressure during induced hypotension. *Anesthesiology* 35:431, 1971.
8. Kirkendall, W. M., Burton, A. C., Epstein, I. H. and Fries, L. D. Recommendations for human blood pressure determination by phycomanometers—Report of a subcommittee of the postgraduate education committee, American Heart Association. *Circulation* 36:980, 1967.
9. Thulemus, O. and Gjores, J. E. Use of Doppler shift detection for determining peripheral arterial blood pressure. *Anology* 22:594, 1971.
10. Kazam, T. M., Gander, M. I., Franklin, D. L. and Ros, J. Blood pressure measurement with Doppler ultrasonic flowmeter. *J. Appl. Physiol.* 30:585, 1971.
11. McLoughlin, G. W., Kirby, E. R., Kemmerer, W. T. and deLemos, R. A. Indirect measurement of blood pressure in infants utilizing Doppler ultrasound. *J. Pediatr.* 9:300, 1971.
12. Blood pressure of adults by age and sex. United

Table II Relation of blood pressure to ECG

| Systolic B P | | Questionable ECG | | Diastolic B P | | Questionable ECG | |
|--------------|-------|------------------|------|---------------|-------|------------------|-------|
| (mm Hg) | No | No | % | (mm Hg) | No | No | % |
| 91 100 | 9 | 1 | 11 | 51 60 | 10 | 4 | 40 |
| 101 110 | 44 | 4 | 9 | 61 70 | 102 | 26 | 26 |
| 111 120 | 149 | 33 | 22 | 71 80 | 321 | 84 | 26 |
| 121 130 | 250 | 61 | 24 | 81 90 | 399 | 125 | 31 |
| 131 140 | 242 | 76 | 31 | 91 100 | 206 | 99 | 48 |
| 141 150 | 198 | 83 | 42 | 101 110 | 54 | 32 | 59 |
| 151 160 | 120 | 56 | 48 | 111 120 | 15 | 9 | 60 |
| 161 170 | 50 | 29 | 58 | 121 130 | 1 | 1 | (100) |
| 171 180 | 20 | 17 | 85 | | | | |
| 181 190 | 13 | 11 | 85 | | | | |
| 191 200 | 9 | 6 | 67 | | | | |
| 201 | 4 | 3 | 75 | | | | |
| Total | 1 108 | 380 | 34.4 | | 1 108 | 380 | 34.4 |

ments was used on 600 participants without failure.

There were 1 108 male participants in the screening program. For each age category the mean weight, number of systolic and diastolic hypertensives and number with abnormal ECGs were tabulated (Table I).

Results

The 1 108 participants had a mean age of 47.5 years. Of the 1 106 participants tested, 19½ per cent had systolic hypertension (> 150 mm Hg) and 6.9 per cent had diastolic hypertension (> 100 mm Hg). Thirty-four per cent had questionable electrocardiograms. The per cent of hypertensives and abnormal ECGs increased with age. Comparison of mean systolic and diastolic BP by age groups with the U.S. population¹² shows the participants to be representative of the population, although both systolic and diastolic tended to be a few mm Hg elevated (Fig. 2). The level of systolic and diastolic pressures were closely related to the percentage of questionable ECGs (Table II and Fig. 3). The proportion of questionable ECGs rose smoothly with the level of blood pressure. No distinct "break point" was evident at any level. A similar rise occurred with age (Table I).

Regression analyses were done between ECG measurements and the following: systolic and diastolic BP, age and ponderal index. Correlations were attempted with P wave amplitude (A) in II, aVL, V₁ and V₂; Q wave A in II, aVL and V₁; Q wave duration (D) in II, aVL and V₁; R wave A in II, aVL and V₁; R wave A + S A in V₁ and V₂; T wave A in II, aVL, V₁, V₂; PR; II, QRS; D; II, QRS frontal plane axis; ST-T angle and QRS-T angle. Regression analyses were also performed between heart rate and age, BP, PR, QRS, D in II, QRS axis, ST-T angle and QRS-T angle. No significant correlations were found.

Discussions and conclusions

Blood pressure is one of the more important prognostic measurements in health evaluation. Manual cuff stethoscope methods have inherent limitations, including observer variation, which may be severe,^{13,14} the requirement for trained personnel and the development of fatigue during prolonged health testing. Automatic methods offer reproducibility and accuracy and are indefatigable. The instrument used in this study has been evaluated for accuracy.^{15,17} This study demonstrates that accurate results may be obtained rapidly by personnel without training in the cuff stethoscope method.

Individual blood pressures are usually not considered hypertensive until a level of > 150 mm Hg systolic or > 100 mm Hg diastolic has been attained. These results showed 19 per cent with systolic and 6.9 per cent with diastolic hypertension. The US figures based on a lower average age are 17 per cent and 7.7 per cent.¹⁷

The differences from the US population mean for each age group are well within the bounds of observer variation^{18, 19} and are thus probably not clinically significant. The small consistent difference noted may be due to the use of the more sensitive ultrasound method or to the activity of subjects attending a meeting.

As expected the blood pressure level rose with age but the body weight did not. The percentage of questionable ECG's is similar to previously reported data on a similar population of dentists.^{14, 15} These data indicate that the population screened at the 1970 Annual Health Evaluation is representative of the US male population with respect to blood pressure. We believe from unpublished data (C. A. Ciceres) that this applies to the ECG as well.

The almost linear relation of the proportion of questionable ECG's to BP was not expected. Another unexpected finding was the lack of correlations of age and BI with the ECG changes they are usually associated with. Review of the age and BP distribution shows the number of extreme cases may be too small to allow this correlation to become clear. Nonetheless the ECG does change with BI but in a non-specific way.

Summary and conclusion

An automatic ultrasound blood pressure instrument was used to measure blood pressure (BI) and a computer was used to analyze ECG's in 1108 male professionals at a voluntary Health Evaluation Program at a professional society meeting. BI's were recorded with the technique recommended by the American Heart Association in an average time of 45 seconds per participant by technicians untrained in the auscultatory method of obtaining BP. Nineteen per cent of the participants had systolic hypertension (> 150 mm Hg) and 7 per

cent had diastolic hypertension (> 100 mm Hg). The mean BP's and percentage of abnormalities were equivalent to the US population. Computer ECG analysis showed 34.4 per cent questionable findings consistent with previous experience. The study showed a strong positive linear relationship between the proportion of questionable ECG's and the systolic and diastolic blood pressure.

The author wishes to thank Mr H. C. Chin and Mr C. Rozen from Roche Medical Electronics who performed the statistical analysis presented here.

REFERENCES

1. Hochberg, H. M., Wehrer, A. I., Abraham, S. and Ciceres, C. A. Automatic ECG analysis. *J. Am. Dent. Assoc.* 3: 644, 1966.
2. Hochberg, H. M., Calistayud, J. H., Wehrer, A. I., Whiteman, J. and Ciceres, C. A. Automatic ECG analysis in rapid mass screening. *Arch. Environ. Health* 15: 190, 1967.
3. Ciceres, C. A. and Hochberg, H. M. The form and use of the computer in diagnosis in the analysis of the electrocardiogram. *Am. Heart J.* 79: 439, 1970.
4. Ciceres, C. A. and Dresfu, I. M. Clinical electrocardiography and computer. New York, 1970. Academic Press.
5. Hochberg, H. M. and Salomon, H. Accuracy of an automated ultrasound blood pressure monitor. *Curr. Ther. Res.* 13: 139, 1971.
6. Sheppard, I. C., Johnson, T. S. and Kirklin, J. W. Controlled study of brachial artery blood pressure measured by a new indirect method. *J. Am. Assoc. Med. Instrum.* 7: 797, 1971.
7. Topper, I. J., Fein, R. M. and Donham, R. T. Automatic ultrasound monitoring of blood pressure during induced hypotension. *Am. J. Physiology* 220: 431, 1971.
8. Kirkendall, W. M., Burton, A. C., Epstein, F. H. and Fries, E. D. Recommendations for human blood pressure determination by phycomanometers—Report of a subcommittee of the postgraduate education committee. American Heart Association. *Circulation* 36: 980, 1967.
9. Thulesius, O. and Gjores, J. E. Use of Doppler shift detection for determining peripheral arterial blood pressure. *Am. J. Physiology* 221: 594, 1971.
10. Kazam, T. M., Gander, M. P., Franklin, D. L. and Ross, J. Blood pressure measurement with Doppler ultrasonic flowmeter. *J. Appl. Physiology* 30: 585, 1971.
11. McLaughlin, G. W., Kirby, R. R., Kemmerer, W. T. and deLemos, R. A. Indirect measurement of blood pressure in infants utilizing Doppler ultrasound. *J. Pediatr.* 9: 300, 1971.
12. Blood pressure of adults by age and sex. United

Table II Relation of blood pressure to ECG

| Systolic B P | | Questionable ECG | | Diastolic B P | | Questionable ECG | |
|--------------|-------|------------------|------|---------------|-------|------------------|-------|
| (mm Hg) | No | No | % | (mm Hg) | No | No | % |
| 91 100 | 9 | 1 | 11 | 51 60 | 10 | 4 | 40 |
| 101 110 | 44 | 4 | 9 | 61 70 | 102 | 26 | 26 |
| 111 120 | 149 | 33 | 22 | 71 80 | 321 | 84 | 26 |
| 121 130 | 250 | 61 | 24 | 81 90 | 399 | 175 | 31 |
| 131 140 | 242 | 76 | 31 | 91 100 | 206 | 99 | 48 |
| 141 150 | 198 | 83 | 42 | 101 110 | 54 | 32 | 59 |
| 151 160 | 120 | 56 | 48 | 111 120 | 15 | 9 | 60 |
| 161 170 | 50 | 29 | 58 | 121 130 | 1 | 1 | (100) |
| 171 180 | 20 | 17 | 85 | | | | |
| 181 190 | 13 | 11 | 85 | | | | |
| 191 200 | 9 | 6 | 67 | | | | |
| 201 | 4 | 3 | 75 | | | | |
| Total | 1 108 | 380 | 34.4 | | 1 108 | 380 | 34.4 |

ments was used on 600 participants without failure

There were 1 108 male participants in the screening program. For each age category the mean weight, number of systolic and diastolic hypertensives and number with abnormal ECGs were tabulated (Table I).

Results

The 1 108 participants had a mean age of 47.5 years. Of the 1 108 participants tested, 19½ per cent had systolic hypertension (> 150 mm Hg) and 6.9 per cent had diastolic hypertension (> 100 mm Hg). Thirty-four per cent had questionable electrocardiograms. The per cent of hypertensives and abnormal ECGs increased with age. Comparison of mean systolic and diastolic BP by age groups with the U.S. population¹² shows the participants to be representative of the population, although both systolic and diastolic tended to be a few mm Hg elevated (Fig. 2). The level of systolic and diastolic pressures were closely related to the percentage of questionable ECGs (Table II and Fig. 3). The proportion of questionable ECGs rose smoothly with the level of blood pressure. No distinct 'break point' was evident at any level. A similar rise occurred with age (Table I).

Regression analyses were done between ECG measurements and the following: systolic and diastolic BP, age and ponderal index. Correlations were attempted with P wave amplitude (A) in II, aV₁, V₂ and V₃; Q wave (Q) in II, aV₁ and V₃; Q wave duration (D) in II, aV₁ and V₃; R wave (R) in II, aV₁ and V₃; R/V₃ A + S in V₁ and V₂; T wave (T) in V₁, V₂, V₃; PR II; QRS D II; QRS frontal plane axis; ST T angle and QRST angle. Regression analyses were also performed between heart rate and age, BP, PR, QRS D in II, QRS axis, ST T angle and QRST angle. No significant correlations were found.

Discussions and conclusions

Blood pressure is one of the more important prognostic measurements in health evaluation. Manual cuff stethoscope methods have inherent limitations, including observer variation, which may be severe,¹³ the requirement for trained personnel and the development of fatigue during prolonged health testing. Automatic methods offer reproducibility and accuracy and are indefatigable. The instrument used in this study has been evaluated for accuracy.^{4,7} This study demonstrates that accurate results may be obtained rapidly by personnel without training in the cuff stethoscope method.

ARRHYTHMIAS

DON'T ALWAYS WAIT FOR YOU!

*A new edition offers guidelines
for expert staff action*

New 2nd Edition!

Phibbs

THE CARDIAC ARRHYTHMIAS

This new edition retains the clarity and simplicity of approach which characterized its predecessor while incorporating a totally modern approach for instructing non cardiologists to diagnose and treat arrhythmias. Stressing the importance of diagnosis through both electrocardiograph interpretation and clinical or bedside diagnosis, it begins with a review of the anatomy and physiology of the heart and the basics of electrocardiography. Dr. Phibbs then leads the reader step by step from normal rhythms to simple arrhythmias and finally to the most complex arrhythmias. With a heavy emphasis on accurate diagnosis before treatment, the book interjects three problem solving sections into its progression to test the reader's comprehension of material covered. After thoroughly guiding the reader to proficiency in diagnosis, a final section discusses drugs and techniques such as defibrillation used in treating cardiac arrhythmias.

By BRENDAN PHIBBS MD FACP FACC with a contribution by GORDON A. EWY MD FACC August 1973 2nd edition 206 pages plus FM I VIII 7 x 10 264 illustrations Price \$7.50

Please send me a copy of Phibbs THE CARDIAC ARRHYTHMIAS (#3910) priced at \$7.50 on 30 day approval. At the end of this time I may keep the book and forward payment or return it without charge or obligation. (I save delivery charges by enclosing payment with my order.)

☐ Bill me ☐ Payment enclosed (Same return privilege guaranteed)

NAME _____

ADDRESS _____

CITY _____

STATE _____ ZIP _____

30-day approval good only in the continental US and Canada

MOSBY

TIMES MIRROR

AMJ1273

THE C. V. MOSBY COMPANY
11830 WESTLINE INDUSTRIAL DRIVE
ST. LOUIS, MISSOURI 63141

- States 1960-1962. National Center for Health Statistics. U.S. Department of Health, Education and Welfare. Public Health Service publication No. 1000 Series 11, No. 4. June 1964.
13. King G L. Taking the blood pressure. *JAMA* 209:1902, 1969.
14. Armitage P and Rose G A. The variability of measurements of casual blood pressure I and II. *J Clin Sci* 30:325, 1966.
15. Edertson E and Humerfelt S. The observer variation in the measurement of arterial blood pressure. *Acta Med Scand* 184:145, 1968.

Sentry 75

ventricular inhibited
pulse generator



GENERAL  ELECTRIC

Index to advertisers

Abbott Laboratories

Enduron 10 11
K Lor 1

Astra Pharmaceutical Products, Inc

Nylocaine HCl Intravenous 22 23 24

Avionics Biomedical Division

Exercise Stress Monitoring 14
Model 660 Electrocardiograph Systems 2

Biotronex Laboratory, Inc

Blood Flowmeter 25

Burroughs Wellcome

Cardilate 27
Lanoxin Second Cover

Cooper Laboratories

Kay Ciel 1

CPC International, Inc

Mazola Oil 26

ESB Medcor, Inc

Pacer Check System 6

General Electric

Sentry 75 Pulse Generator 17 18 19 20

Ives Laboratories, Inc

Isordil 8

Marion Laboratories, Inc

Nitro Bid 12

Lever Brothers Company

Pronase Marcamine 1

Marquette Electronics, Inc

Electrocardiography Equipment Third Cover

Purdue Frederick

Cardioquon Fourth Cover

Roche Laboratories

Valium 28 29 30

New circuit efficiencies through the development and adaptation of thick film hybrid circuitry born of aerospace technology almost one-half the normal electrical interconnections are eliminated. The new circuits improve electrical efficiency by 45% (135 micro-watts of power drain vs. 244 micro-watts formerly).



The new micro-circuitry in both the sensing and pacing circuits—plus the rectangular power cells—allow a 25% reduction in the volume of the generator. Thus while generator size is reduced power capacity is increased 50%.

Hermetic sealing of these critical hybrid circuits plus a second overall hermetic sealing of the complete circuitry virtually locks-out

deteriorative moisture The metal hermetic containers also provide additional protection from electromagnetic interference (EMI).

In addition patient defibrillation can be conducted without damaging the generator

Implant procedures are not altered because the Sentry 75 is compatible with all existing GE electrodes which feature simple self sealing push pull connectors. It is also adaptable to the electrodes of other manufacturers.

The pulse generator is provided as a bipolar system which at implant can be converted to unipolar pacing if desired. This is accomplished by simply removing the insulating label covering the metallic plate electrode on the generator's surface and attaching it to a unipolar electrode or to a bipolar electrode adapted for unipolar pacing.

Choice of rates. Two models of the Sentry 75 pulse generator are presently available with two additional models scheduled for subsequent availability. Models differ only in pacing and hysteresis rates.

| Model | Pace Rate | Hysteresis (Escape Rate) |
|-----------|-----------|--------------------------|
| A2075A/AD | 73 | 63 |
| A2075B/BD | 80 | 70 |
| A2075C/CD | 85 | 80 |
| A2075D/DD | 73 | 71 |

A 14 sec. delay of 1974

GENERAL  ELECTRIC

Sentry 75... one generator that meets long-term pacing demands

The Sentry 75 ventricular inhibited pulse generator maximizes the potentials of a proven power source to deliver extended pacing life to most patients regardless of threshold value

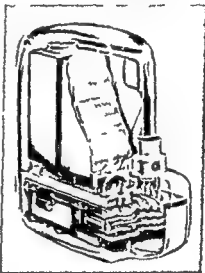
It is General Electric's positive response to the medical profession's demand for a simple extended life pacing system to serve the large majority of patients

The Sentry 75 presents no alterations in implant technique or patient management. It is compatible with all GE electrodes and the long reliable service life can be monitored easily by both patient and physician

The Sentry 75 is powered by the industry's first mercury/zinc cells specifically designed for implantable pulse generator applications. A complete redesign of both this power cell and circuitry now make possible a pulse generator which can be expected to deliver a 4 to 6 year life with an assurance factor greater than generators using normal mercury/zinc batteries

Your GE representative can provide you with information regarding patient testing, warranties, and availability in your area

Optimized power source capability



Redesigned power cells The mercury/zinc battery is a long proven power source familiar to the medical profession. Its chief liability has not been in the chemical composition but in its construction. GE design aims were to maximize power utilization and reduce internal losses to produce a power cell capable of far longer and more reliable service life.

Note the new GE power cell is rectangular. It displaces the same volume as 2 circular batteries yet contains 50% greater capacity (3000 mA hours vs. 2000) than 2 circular batteries operating in parallel. Two of these high capacity cells connected in series

power the pulse generator. Internal discharge is a problem with previous batteries resulting from shorting electrodes. This has been virtually eliminated by 11 wraps of a non-degradable separator. These multi wraps replace single wrap separator presently used in mercury zinc batteries.

The power cell case is soldered nickel to prevent case-to-electrolyte reaction, another potential source of self-discharge.

Welded internal contacts and electrodes preclude the possibility of open contacts, a design shortcoming with ordinary pressure contacts.

The negative terminal is isolated from the cap through a hermetic ceramic-to-metal seal, and the cap is welded to the top of the cell to prevent electrolyte seepage across 2 terminals. And as one last safety precaution, the post-pressure vent is located at the end of the cell, remote from the terminals.

Experimental and laboratory reports

Electrophysiological evaluation of disopyramide in man

Mark E Josephson MD

Anthony R Caracta MD

Sun H Lau MD

John J Gallagher MD

Anthony N Damato MD

Staten Island NY

Disopyramide (Norpace)* is a new antiarrhythmic agent available in Europe (Rythmodin) with a reported antiarrhythmic spectrum similar to that of quinidine. It has been used orally and parenterally and has been found useful in the management of paroxysmal arrhythmias and extrasystoles of ventricular or supraventricular origin.^{1,2} This study was undertaken to analyze the effects of intravenously administered disopyramide on the refractory periods (RP) of the atrium (A), A-V node (A-VN) and His-Purkinje system (HPS) as well as on conduction through the A-VN and HPS in man.

Methods and materials

Twelve patients underwent right heart catheterization in the nonsedated postabsorptive state after informed consent was obtained. Clinical data are listed in Table I. Five patients had arrhythmias and/or conduction disturbances while the others were studied during the course of catheterization for other reasons.

A quadripolar electrode catheter was

inserted percutaneously into an antecubital vein and positioned fluoroscopically against the lateral wall of the right atrium. The two distal electrodes were used to pace the atrium while the proximal pair recorded a high right atrial (HRA) electrogram. His bundle electrograms (HBE) were obtained according to methods previously described.^{3,4} Three or four surface electrocardiographic leads were simultaneously recorded (I, II, III, +I, -I). Each patient was paced at several basic cycle lengths for 30 to 60 seconds during which time records were made of A-V nodal and His-Purkinje conduction time (vide infra). Refractory periods were determined by the extra-stimulus method⁵ using a specially built programmed digital stimulator which delivered rectangular impulses of 1 msec duration at twice diastolic threshold. Progressively more premature impulses were delivered after every eighth spontaneous or paced beat until the effective refractory period (ERP) of the atrium was reached. Refractory period studies were carried out at paced atrial cycle lengths in 11 patients

From the Cardiac Electrophysiology Laboratory, St. Elizabeth's Hospital, Staten Island, NY.
This work was supported in part by the Federal Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, MD.
Received for publication July 15, 1973.
Reprint requests to Mark E. Josephson, MD, Cardiac Electrophysiology Laboratory, St. Elizabeth's Hospital, 1000
Hudson Avenue, Staten Island, NY 10310.
Kindly published by G.D. Seltzer and C. Clinego III.

Patient management

The Sentry 75 pulse generator has a built in service life indicator that is simple and easy to use. The pulse generator's pacing rate is highly dependent upon battery voltage; thus as battery voltage begins to fall near end of life, pacing rate will decrease. This change in rate can be easily and accurately monitored when a GE patient sensor and rate interval computer are used. The latter is a desk top unit which, used in conjunction with the patient sensor, enables the physician to check the patient's pacemaker over the phone between regular office visits. The patient can also be instructed to check his own rate with the sensor.



Clinical evaluation

Laboratory tests of the Sentry 75 components were conducted for 2 years before any implants in humans were undertaken. Clinical evaluations began on a worldwide basis in November 1972. Physicians representing more than 15 major hospitals and medical centers are participating in the program.

General Electric
Medical Systems Milwaukee
Toronto Liege Madrid



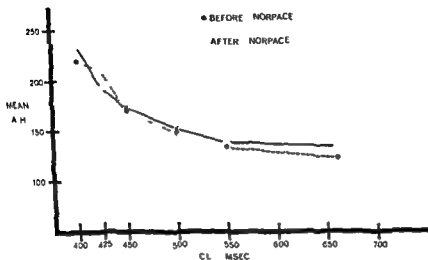


Fig 1 This graph demonstrates the effects of disopyramide on AV nodal conduction at different paced rates. Mean A-H is plotted on the ordinate and paced cycle length on the abscissa. It can readily be seen that there is no significant difference in response after disopyramide (Norpace).

Table III Effect of disopyramide on the ERP of the atrium*

| Patient No | Cycle length at which study done | ERP before Norpace | ERI after Norpace |
|-----------------------------------|----------------------------------|--------------------|-------------------|
| 1 | 500 | 290 | 290 |
| 2 | NSR† | 340 | 340 |
| 3 | NSR | 200 | 260 |
| | 550 | 200 | 210 |
| 4 | 600 | 230 | 240 |
| 5 | NSR | 340 | 320 |
| | 860 | 280 | 280 |
| | 600 | 210 | 280 |
| | 500 | 280 | 270 |
| 6 | 600 | 290 | 280 |
| 7 | 850 | 290 | 280 |
| | 800 | 260 | 275 |
| 8 | NSR | 270 | 260 |
| | 700 | 260 | 210 |
| | 600 | 210 | 270 |
| 9 | 750 | 245 | 280 |
| | 600 | 235 | 260 |
| 10 | 700 | 240 | 270 |
| | = 1 mg/kg | | |
| | 600 | 45 | 270 |
| | 700 | 240 | 265 |
| | = 1.5 mg/kg | | |
| | 600 | 240 | 265 |
| 11 | NSR | 60 | 260 |
| 12 | NSR | 310 | 260 |
| | 600 | 255 | 290 |
| Mean ERI ± standard error of mean | | 256 ± 6 | 269 ± 4 |
| Net change = 13 | | | |
| P value 0.05 | | | |

* 1 msec
† NSR = no mal
hythm

Table I Clinical data

| Patient No | Age | Sex | Weight (kg) | Dose of disopyramide (mg) | Heart disease | Medication |
|------------|-----|-----|-------------|---------------------------|---------------------|-------------|
| 1 | 66 | M | 70 | 140 | ASHD | None |
| 2 | 41 | M | 80 | 120 | None | None |
| 3 | 55 | M | 60 | 90 | Mitotonic dystrophy | None |
| 4 | 52 | M | 60 | 60 | ASHD | Off digoxin |
| 5 | 54 | M | 85 | 135 | ASHD | None |
| 6 | 40 | M | 80 | 80 | None | None |
| 7 | 53 | M | 70 | 140 | ASHD | None |
| 8 | 36 | M | 85 | 85 | ASHD | None |
| 9 | 18 | M | 80 | 80 | Mitotonic dystrophy | None |
| 10 | 74 | M | 70 | 70 | ASHD | None |
| | | | | 100 | | |
| 11 | 67 | M | 90 | 180 | ASHD | None |
| 12 | 45 | M | 60 | 90 | ASHD | None |

Table II Effects of disopyramide on A-V nodal conduction at paced rates*

| Patient No | 660 | | 550 | | 500 | | 450 | | 405 | | 400 | |
|------------|----------|----------|---------|----------|----------|----------|----------|----------|----------|----------|---------|----------|
| | B | A | B | A | B | A | B | A | B | A | B | A |
| 1 | 110 | 100 | 110 | 120 | 130 | 140 | 150 | 150 | 200 | 200 | 200 | W |
| 2 | 110 | 130 | 130 | 145 | W | W | — | — | — | — | — | — |
| 3 | — | — | 140 | 150 | 150 | 150 | 150 | 150 | 210 | 225 | 0 | 210 |
| 4 | 100 | 95 | 110 | 100 | 115 | 110 | 120 | 115 | W | 120 | W | 15 |
| 5 | 110 | 110 | 135 | 110 | 145 | 120 | 150 | 150 | 200 | 115 | W | 155 |
| 6 | 90 | 135 | 115 | 160 | 100 | 170 | 210 | 280 | 215 | W | — | — |
| 7 | 90 | 100 | 110 | 110 | — | — | — | — | — | — | — | — |
| 8 | 130 | 155 | 175 | 190 | 210 | 00 | W | 220 | — | W | — | — |
| 9 | 200 | 200 | W† | 300 | — | W | — | — | — | — | — | — |
| 10 | 175 | 160 | 180 | 170 | 205 | 190 | 215 | 195 | 245 | 225 | W | W |
| 11 | 110 | 90 | 100 | 105 | 100 | 105 | 140 | 115 | 235 | 125 | W | 245 |
| 12 | — | — | — | — | 145 | 165 | — | — | 170 | 155 | 200 | 200 |
| Mean A-H | 123 ± 12 | 134 ± 16 | 133 ± 8 | 136 ± 10 | 151 ± 12 | 159 ± 12 | 172 ± 13 | 174 ± 21 | 210 ± 16 | 210 ± 16 | 210 ± 0 | 235 ± 35 |
| + SEM | | | | | | | | | | | | |
| Net change | + 11 | | + 3 | | + 1 | | + 2 | | + 17 | | + 15 | |
| % change | 9% | | 23% | | 7% | | 1% | | 81% | | 67% | |
| P value | 2 | | > 5 | | > 5 | | > 5 | | > 5 | | > 5 | |

*Values are expressed in msec
†W = Wenckebach

to avoid the effect of changing cycle length on refractoriness. In 7 patients refractory periods were also studied during normal sinus rhythm.

After control values were obtained each patient was given 1 to 2 mg per kilogram of body weight of disopyramide intravenously over a 5 to 10 minute period. One

patient (No. 10) was studied at two dose levels. Studies of conduction and refractory periods were then repeated. Blood pressure was frequently measured throughout the study with a sphygmomanometer. Careful attention was paid to note any clinical signs of myocardial depression.

Time lines were generated at 10 and 100

Table IV Effects of disopyramide on the ERP of the AVN*

| Patient No | CL ₁ | B | I |
|------------|------------------|----------|----------|
| 1 | 500 | 330 | 360 |
| 2 | NSR ₁ | 350 | 340 |
| | 100 | 450 | 400 |
| 4 | NSR | 300 | 260 |
| | 600 | 310 | 210 |
| 5 | 500 | 310 | 270 |
| 6 | 600 | 380 | 280 |
| 8 | NSR | 330 | 315 |
| | 100 | 310 | 360 |
| | 600 | 410 | 390 |
| 9 | 750 | 385 | 390 |
| | 600 | 460 | 450 |
| 10 | 100 [†] | 330 | 330 |
| | 600 [†] | 350 | 350 |
| | 100 [‡] | 330 | 330 |
| | 600 [‡] | 350 | 340 |
| 12 | 600 | 295 | 295 |
| | Mean ± SEM | 366 ± 13 | 344 ± 14 |
| | Net change | -22 | |
| | P value | 05 | |

*val msec.

† 0 mg disopy am da.

‡ 100 mg disopy mid

Abb CL = CL₁ length NSR = normal rhythm.

longest H₁H₂ interval at which H₂ fails to depolarize the ventricles

The relative refractory period (RRP) of the HIS is defined as the longest H₁H₂ interval at which H₂ conducts to the ventricles with a longer H₁V interval than that of the basic drive beat or with a QRS of aberrant configuration. This definition presupposes that the HPS functions as a single unit. Although it is recognized that the HPS is a trifascicular system in the absence of multiple recording sites along individual fascicles it is impossible to precisely measure the ERP versus the RRP of any given fascicle. Thus for the purposes of this study it was elected to consider the HPS as a single functioning unit and the RRP as so defined.

Results

Spontaneous sinus cycle length was unchanged or insignificantly (< 100 msec) decreased in 11 patients but in 3 patients (Nos 3, 4 and 5) it was shortened by 120 to 160 msec.

AV nodal conduction Disopyramide variably and insignificantly affected AV nodal conduction during sinus rhythm increasing it in five decreasing it in two and not changing it in the remaining five patients. Table II summarizes the effects of disopyramide on AV nodal conduction as measured by AH interval at several paced rates. At any paced atrial rate AV nodal conduction was not significantly altered by disopyramide (Fig 1). However it should be noted that in 5 patients (Nos 4, 5, 8, 9 and 11) AV nodal Wenckebach occurred at higher paced rates than during control. A sixth patient (No 10) showed enhancement of AV nodal conduction although Wenckebach developed at the same paced rate as it did during control studies. Thus several patients showed enhancement of AV nodal conduction during atrial pacing compared to control.

HIS Purkinje conduction Control HV values were normal in 9 patients (35 to 55 msec) and abnormal in 3 (Nos 3, 11, 12—60 to 62 msec). In 10 of 12 patients

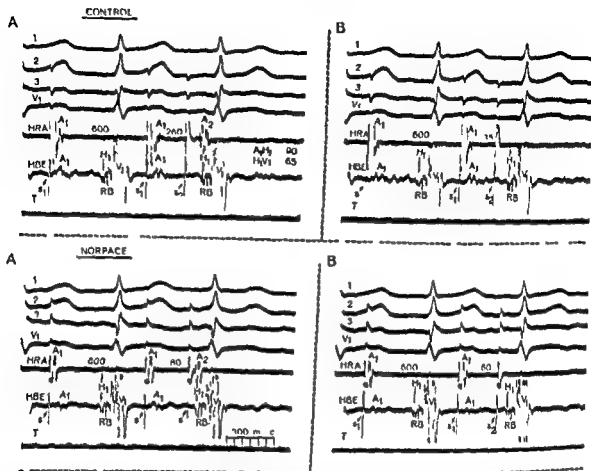


Fig 2 A and B Effects of disopyramide (Norpace) on the ERP of the atrium. Panels A and B in the control and after disopyramide represent the shortest A₁A₂ (panel A) and S₁S₂ at which S₂ fails to capture the atrium—the 1:1 RI of the atrium (panel B). In this patient at a paced cycle length of 600 msec the ERP of the atrium is increased 25 msec (235 to 260 msec) after disopyramide.

msec. All data were stored on magnetic tape* and later retrieved at paper speeds of 150 mm per second†.

A thirteenth patient not listed in Table I who did not undergo cardiac catheterization was given a 2 mg per kilogram of body weight bolus of disopyramide during an episode of supraventricular tachycardia and electrocardiographic changes were recorded.

Statistical data were calculated using the Student t test for paired data. In evaluation of refractory periods statistical analysis was only applied to those determinations obtained during paced atrial cycle lengths. This eliminates the effects of changing cycle length or refractoriness.

Definition of terms

The AH (atrium to His) interval as measured in the His bundle electrogram recording was taken as a measure of AV nodal conduction time (nl 60 to 140 msec).

The HV (His to ventricle) interval was taken as a measure of His Purkinje conduction time (nl 30 to 55 msec).

The effective refractory period (ERP) of the atrium is defined as the longest S₁S₂ that does not result in atrial depolarization.

The ERP of the AVN is defined as the longest A₁A₂ interval at which A₂ depolarizes the atrium but does not propagate to the HPS.

The functional refractory period (FRP) of the AVN is defined as the shortest H₁H₂ interval resulting from any two consecutive propagated atrial depolarizations.

The ERP of the HPS is defined as the

*North American Phillips Shelton Conn
†Electronics for Medicine White Plains N.Y.

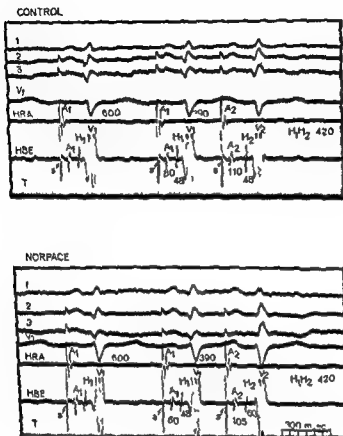


Fig 4 Effect of disopyramide (Norpace) on the RRP of the HPS. This patient while being paced at a cycle length of 600 msec showed no aberration or H1H2 prolongation at an H1H2 of 420 msec before disopyramide. After disopyramide at the same H1H2 of 420 msec the H1H1 is prolonged to 60 msec and aberration in the form of incomplete LBBB is present. Thus in this patient the RRP of the HPS was prolonged by disopyramide.

mined since right heart catheterization was not performed. In this patient disopyramide slowed the ventricular response from 100 to 120 (Fig 7). Continued oral disopyramide 100 mg QID for two days converted the arrhythmia to sinus rhythm.

Side effects. The only side effect noted was dry mouth present in two patients. There was no significant hypotension, myocardial depression or conduction disturbances noted.

Discussion

Disopyramide is a new antiarrhythmic agent which shares many clinical properties with quinidine even though its chemical structure is very different. It has received many trials in Europe and apparently has its major use in the treatment of supra-

ventricular and ventricular extrasystoles and in the prophylaxis of paroxysmal arrhythmias.¹⁻⁶ There are conflicting data on its usefulness in chronic atrial fibrillation.⁶ A few studies have shown it to be effective in the prevention of arrhythmias after acute myocardial infarction.⁷ It may be given orally in doses of 100 to 200 mg QID or may be given intravenously at a dose of 1 to 2 mg per kilogram of body weight. Its action is relatively rapid when given intravenously.

Little is known about the electrophysiological properties of disopyramide in man. Several studies¹¹⁻¹³ in dogs found dose related increases of the A-V functional RP and A-V conduction time. There is also some evidence that disopyramide may slow His-Purkinje conduction and diminish

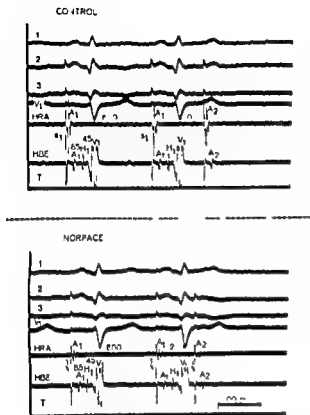


Fig 3 Effects of disopyramide (Norpace) on the LRP of the AVN. This patient while being paced at a cycle length of 600 msec reached the LRP of the AVN at an $A_1 A_2$ of 370 msec during control studies. After disopyramide block in the node first appears at an $A_1 A_2$ of 270 msec. Thus the LRP of the AVN was shortened 100 msec.

disopyramide had no effect on H V intervals at sinus rhythm or at various paced atrial rates. In two patients (Nos 3 and 5) the H V interval increased by 5 and 10 msec (50 to 55 and 60 to 70 msec) after 1 mg per kilogram of body weight of disopyramide.

Refractory studies Although refractory periods were determined at sinus rhythm as well as during paced rates in most patients only those values obtained during paced cycle lengths were considered valid due to the known effects of changing cycle lengths on refractoriness. Therefore all calculations of mean and P values were derived from the results of only those studies done at paced cycle lengths.

ERP of the atrium Table III summarizes the effects of disopyramide on the ERP of the atrium. The ERP of the atrium was increased by 13 msec. This is a small but statistically significant change ($p = .005$).

In only three patients did the ERP decrease after disopyramide. Fig 2 is from patient No 9 in whom disopyramide lengthened the ERP of the atrium from 73 to 260 msec.

ERP of the AVN The ERP of the AVN could be measured in only nine patients. In the remaining 3 patients the ERP of the atrium was reached before that of the AVN. The results are summarized in Table IV. On the average the ERP of the AVN was shortened 22 msec after disopyramide. This is significant to a P value of .05. Fig 3 demonstrates a 100 msec shortening of the ERP of the AVN after disopyramide.

LRP of the AVN The LRP of the AVN was not significantly changed by disopyramide in spite of the fact that the ERP of the AVN was shortened.

ERP and RRP of the HPS No patient developed block within the HPS during paced rates; therefore no comment can be made on disopyramide's effect on this parameter. One patient did develop block in the HPS during NSR and in this patient disopyramide shortened the ERP by 20 msec. The meaning of this is not clear since this patient had changing atrial cycle lengths at NSR both before and after disopyramide and thus may have altered the refractoriness of the HPS.

Disopyramide prolonged the RRP of the HPS in two patients. In one patient the RRP increased by 20 msec at two paced rates after disopyramide. In the other patient incomplete LBBB developed after disopyramide where no aberration was present prior to its administration. This is shown in Fig 4.

Runs of supraventricular tachycardia were noted in two patients (Nos 12, 13). One patient (No 12) developed repeated episodes of A V nodal reentrant type paroxysmal atrial tachycardia (PAT) in response to electrically induced atrial depolarizations. In this patient disopyramide shortened the critical A V nodal delay required to produce PAT from 190 to 175 msec. However the R R interval of the reentrant rhythm was unchanged. The second patient (No 13) spontaneously developed a supraventricular tachycardia the mechanism of which could not be deter-

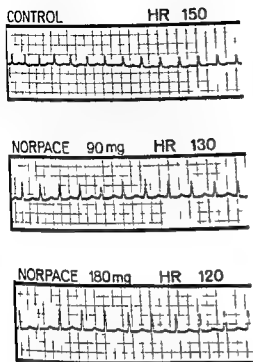


Fig 5 Effect of disopyramide (Norpace) on supraventricular tachycardia. This patient developed a supraventricular tachycardia with a ventricular response of 150 beats per minute. After 180 mg of disopyramide the ventricular rate had gradually decreased to 120 beats per minute. No discrete P waves are seen.

nodal refractoriness. Alternatively, since none of the studies in dogs utilized His bundle electrograms the site of increased refractoriness could not be accurately determined. Our studies demonstrated that in the small number of cases where it could be measured the RRP of the HIS was increased, thus the reported increase in atrioventricular refractoriness may have been related to disopyramide's effect on the HPS, not the AVN. This effect of shortening the ERP of the AVN may have clinical relevance since an increase in ventricular response during treatment of atrial flutter and fibrillation has been seen.^{2,5}

Although we did not correlate electrophysiological effects of disopyramide with plasma levels, the doses used are considered therapeutic.³ Oral doses as low as 100 mg per day have been effective in the treatment of arrhythmias.³ Intravenously administered disopyramide has a relatively rapid onset of action and a half life of

14 to 2 hours.¹² Since our studies took approximately 30 minutes we felt therapeutic levels of disopyramide were present. In no case was the severe myocardial depression¹³ and hypotension¹³ seen in dogs noted.

Conclusion

The results presented above suggest that disopyramide

- (1) has no significant effect on sinus automaticity
- (2) has no significant effect on conduction through the AVN or HPS
- (3) produces a small but probably clinically insignificant increase in the ERP of the atrium
- (4) shortens the ERP of the AVN without significantly affecting the FRP of the AVN
- (5) prolongs the RRP of the HPS
- (6) has no significant side effects at the doses used in the present study
- (7) has an unclear mechanism of antiarrhythmic action

The author wishes to acknowledge the assistance of Audrey Pedersen, Mary Vecchione, Michael Moretti, Anne Mazzella and Kenneth Donohue.

REFERENCES

- 1 Colonna D, Sorocque M, Tep-Tho and Calisti G. *Coeur Med Interne* 455 1968. Translated by I. Lutz.
- 2 Desruelles J, Gerard A, Ducatillon P and Herbauts A. Nos premiers essais cliniques de disopyramide (H 3792) dans les troubles du rythme cardiaque. *Therapie* 22 937 1967.
- 3 Katz M J, Meyer C E, El Etr A and Slodis S J. Clinical evaluation of a new antiarrhythmic agent SC 031. *Curr Ther Res* 343 1963.
- 4 Mokler C M and Van Arman C G. Pharmacology of a new antiarrhythmic agent Disopyramide (2-pyridyl) butylamide 55C 7031. *J Pharmacol Exp Ther* 136 114 1962.
- 5 Searle G D. Norpace (disopyramide phosphate) SC 031 (phosphate). Investigation Brochure 1971.
- 6 Graner J. Un nouvel antiarrhythmique 6 disopyramide. *Pres Med* 6 1605 1968. Translated by I. Lutz.
- 7 Schelegel B J, Lau S H, Helfant R H, Berkowitz W D, Stein E and Damato A N. Catheter technique for recording His bundle activity in man. *Circulation* 39:113 1969.
- 8 Damato A N, Lau S H, Helfant R H, Stein E, Berkowitz W D and Cohen S I. Study of atrioventricular conduction in man.

Table V Effect of disopyramide on the IRP of the AVN*

| Patient No | CL† | Before | After |
|------------|-----|----------|----------|
| 1 | 500 | 430 | 450 |
| 2 | NSR | 490 | 440 |
| | 700 | 560 | 480 |
| 3 | NSR | 380 | 400 |
| | 550 | 360 | 380 |
| 4 | NSR | 390 | 415 |
| | 600 | 420 | 360 |
| 5 | NSR | 430 | 410 |
| | 860 | 400 | 420 |
| 6 | 600 | 495 | 495 |
| 7 | 850 | 420 | 435 |
| | 800 | 415 | 430 |
| 8 | NSR | 450 | 465 |
| | 700 | 470 | 475 |
| | 600 | 470 | 480 |
| 9 | 750 | 560 | 550 |
| | 600 | 550 | 510 |
| 10 | 400 | 445 | 445 |
| | 600 | 435 | 435 |
| | 700 | 445 | 430 |
| | 600 | 435 | 425 |
| 11 | NSR | 410 | 405 |
| 12 | NSR | 440 | 430 |
| | 600 | 415 | 430 |
| Mean ± SEM | | 447 ± 14 | 442 ± 11 |
| Net change | | -5 | |
| t value | | > 5 | |

*Values in msec

†Abbreviations: CL = cycle length; NSR = normal sinus rhythm

phase 4 depolarization of the HPS in dogs.⁶ These studies all used doses that were above what is considered therapeutically safe in man. Studies in man have shown variable ECG changes including prolonged PR and QT interval, widened QRS, I and U wave changes in 2 to 25 per cent of cases.¹⁻⁵ None of these previous reports separated the effects of disopyramide on the AVN and HPS. Using the present methods it has been possible to determine the electrophysiological properties of drugs in man.^{14,16} The results of this study reveal that disopyramide caused no significant change in A-V nodal conduction time during sinus rhythm or at paced rates up to 150. Only two patients show an increase in His-Purkinje conduction time which was minimal (5 and 10 msec) and two patients (Nos 9 and 12) whose control H-V intervals were already prolonged had no further depression of His-Purkinje conduction. Thus, unlike

procainamide^{16,17} and quinidine^{11,12} which depress AVN and HPS conduction, disopyramide does not and might therefore be indicated in the treatment of arrhythmias which are associated with some degree of atrioventricular block.

Our study confirms data from studies in dogs that the ERP of the atrium was increased. Although our result was statistically highly significant ($p = 0.05$) the increase of 13 msec is probably not of clinical significance. This might explain its variable success in the conversion of atrial fibrillation to NSR.

The fact that disopyramide shortened the ERP of the AVN in man came as a surprise in view of all animal work which showed a dose-related increase in A-V nodal refractoriness.^{11,12} There are two possible explanations for this discrepancy. Disopyramide has anticholinergic properties⁴ which may have caused a decrease in A-V

Are patients with essential hypertension and low renin protected against stroke and heart attack?

Roland Stroobandt MD

Robert Fagard MD

Antoon A. P. C. Amery MD

Leuven Belgium

Brunner and colleagues¹ reported recently on 219 patients with essential non malignant hypertension without heart failure. Their plasma renin activity was related to daily sodium excretion and compared to a normogram from 52 normal volunteers studied over the same continuous range of sodium balance. On the basis of this plasma renin level patients were divided in three groups: patients with abnormally low plasma renin level (27 per cent), patients with normal plasma renin level (57 per cent) and patients with abnormally high plasma renin level (16 per cent). Compared to the first two groups the high renin group had higher diastolic blood pressure and more cardiovascular complications of high blood pressure (14 per cent). Apart from the renin level the groups with low and normal renin level did not differ from each other according to other known risk factors for cardiovascular disease including diastolic blood pressure. Nevertheless strokes and heart attacks were not noted in the low renin group and were present in 11 per cent of the patients with normal renin levels. The

authors conclude that plasma renin activity emerges as a potential risk factor for patients with essential hypertension and even suggest that those antihypertensive agents that can correct renin abnormalities could produce a more satisfactory blood pressure response.

The purpose of the present paper is first to present the results of an analysis of these criteria in our patients with essential hypertension in an attempt to test the Laragh hypothesis and second to comment on both studies.

Selection of patients

The last 59 patients hospitalized in our department and discharged with the final diagnosis of essential hypertension were included in this retrospective study in respect of severity and complications of hypertension.

Methodology

Blood for renin determinations was withdrawn in basal conditions i.e. after at least four weeks interruption of diuretics antihypertensive agents and oral con-

From the Department of Medicine I

Received for publication July 15, 1973

Reprint requests to Prof. A. Amery, MD, I

Belgium

University Hospital St. Rafael, Leuven, Belgium

University Hospital St. Rafael, Kapucijnenvoer 35, 3000 Leuven

- using electrode catheter recordings of His bundle activity. *Circulation* 39:287 1969
- 9 Krayer O, Mindok J J and Mendez C. Studies on veratrum alkaloids. VI. The action of epinephrine and of veratramine on the functional refractory period of the atrioventricular transmission in the heart lung preparation of the dog. *J Pharmacol Exp Ther* 107:412 1951
 - 10 Goldreyer B N and Dimato A N. The essential role of atrioventricular conduction delay in the initiation of paroxysmal supraventricular tachycardia. *Circulation* 43:679 1971
 - 11 Dein R R and Terkun D M. Effects of disopyramide on the AV conduction system. *Arch Int Pharmacodyn Ther* 190:183 1971
 - 12 Ranney R I, Dein R R, Karim A and Radziulowski I M. Disopyramide phosphate. Pharmacokinetic and pharmacologic relationships of a new antiarrhythmic agent. *Arch Int Pharmacodyn Ther* 191:167 1971
 - 13 Sekiya A and Aouchim William I M. Comparison of antiarrhythmic actions and effect and intracellular cardiac potentials of pronethalol, disopyramide and quinidine. *Br J Pharmacol* 21:473 1963
 - 14 Dimato A N and Lau S H. Clinical value of the electrogram of the conduction system. *From Cardiovasc Dis* 13 (2):119 1970
 - 15 Josephson M I, Carietta A R, Dimato A N, Gilligher J J and Lau S H. Effects of lidocaine on refractory periods in man. *Am Heart J* 84:748 1972
 - 16 Josephson M I, Carietta A I, Ricciutti M A, Lau S H and Dimato A N. Electrophysiologic properties of procaine amide in man as correlated with plasma level. *Circulation* 46 (Suppl 11):171 1972
 - 17 Rosen K M, Berkowitz W D, Lau S H and Dimato A N. The effects of procaine amide on atrioventricular conduction in man. (Abstr.) *Circulation* 39 and 40 (Suppl 3):October 1969
 - 18 Lau S H, Virghe P J, Bobb C A and Dimato A N. Depression of intra atrial and His Purkinje conduction by quinidine toxicity in dogs. *Circulation* 43 (Suppl 11):189 1971
 - 19 Mathur P P. Cardiovascular effects of a newer antiarrhythmic agent-disopyramide phosphate. *Am Heart J* 84:664 1972

F 1966
V 100 6

Table III Biochemical data in patients with essential hypertension divided into three groups according to the plasma renin concentration

| | Low renin | Middle renin | High renin |
|--|----------------|----------------|----------------|
| | 20 | 19 | 20 |
| Total number of patients | 18 — 65 | 8 — 134 | 14 — 49 |
| Plasma renin concentration range (Skinner ² U/ml) ($\bar{x} \pm S D$) | 4.5 \pm 1.5 | 10.5 \pm 1.9 | 22.3 \pm 8.5 |
| Plasma cholesterol ($\bar{x} \pm S D$) (mg %) | 238 \pm 40 | 225 \pm 37 | 254 \pm 56 |
| Plasma lipid ($\bar{x} \pm S D$) (mg %) | 630 \pm 161 | 511 \pm 113 | 643 \pm 117 |
| Serum creatinine ($\bar{x} \pm S D$) (mg %) | 1.21 \pm 0.4 | 1.15 \pm 0.4 | 1.14 \pm 0.3 |
| Creatinine clearance ($\bar{x} \pm S D$) (ml/min) | 79 \pm 28 | 83 \pm 22 | 93 \pm 29 |

Table IV Some cardiovascular complications in patients with essential hypertension divided into three groups according to the plasma renin concentration

| | Low renin | Middle renin | High renin |
|-----------------------------|-----------|--------------|------------|
| | 20 | 19 | 20 |
| Number of patients at risk | 2 | 1 | 1† |
| Patients with heart failure | 1 | 2 | 1† |
| Patients with stroke | 1 | 1 | 3† |
| Patients with heart attack | | | |

*One patient had both heart failure and heart attack.

†One patient had all three complications.

using the method of Skinner² with minor modifications. With this method normal values range from 8 to 25 units per ml. On the basis of the renin determinations the 59 patients were divided in three tertiles containing 20 or 19 patients each; the three groups thus constituted were compared with each other.

Other analysis of serum, plasma or urine were made with the standard Auto Analyzer technique except for urinary sodium which was measured by internal standard flame photometry.

The blood pressure was measured at the time of blood sampling for renin using a standard sphygmomanometer and taking Phase 5 as diastolic blood pressure.

Results

According to the described techniques one third of the patients ($n = 20$) had a basal plasma renin concentration below 7 units per ml; the plasma renin level of the

middle tertile (19 patients) ranged from 7 to 13.9; the renin level of the upper tertile ($n = 20$) was 14 units or above.

Before studying the incidence of stroke and heart attack in these three groups we analyzed first if the three groups were comparable concerning other known risk factors for cerebrovascular and coronary heart disease. As shown in Tables I, II and III no significant differences were found between the three groups in the incidence of the different mentioned factors.

Table IV and Fig. 1 show the incidence of stroke, heart failure and heart attack in these three groups. No significant differences were found.

In the study of Brunner and colleagues¹ patients with heart failure and malignant hypertension were not entered. If we withdraw the patients with heart failure from our study we retain one patient with stroke and one patient with heart attack.

Table I Data from history in patients with essential hypertension divided into 3 groups according to the plasma renin concentration

| | Low renin | Middle renin | High renin |
|--|-----------------|-----------------|-----------------|
| Total number of patients | 20 | 19 | 20 |
| Sex | | | |
| Male | 11 | 10 | 6 |
| Female | 9 | 9 | 14 |
| Age ($\bar{x} \pm S D$) yr | 46.2 \pm 15.3 | 45.3 \pm 10.6 | 46.7 \pm 14.5 |
| Number with positive family history for stroke, hypertension or heart attack | 14 | 9 | 15 |
| Number of smoking | | | |
| No cigarettes | 11 | 7 | 14 |
| Less than 10 cigarettes a day | 2 | 3 | 2 |
| 10 or more cigarettes a day | 7 | 8 | 4 |
| Unknown | 0 | 1 | 0 |
| Known duration of hypertension ($\bar{x} \pm S D$) yr | 4.7 \pm 5.9 | 7.4 \pm 1.9 | 10.0 \pm 8.6 |

Table II Physical examination in patients with essential hypertension divided into three groups according to the plasma renin concentration

| | Low renin | Middle renin | High renin |
|---|-----------------|----------------|----------------|
| Total number of patients | 20 | 19 | 20 |
| Height ($\bar{x} \pm S D$) (M) | 1.64 \pm 0.1 | 1.67 \pm 0.1 | 1.61 \pm 0.1 |
| Weight ($\bar{x} \pm S D$) (kg) | 69.3 \pm 10.1 | 75.0 \pm 14 | 71.7 \pm 14 |
| Systolic blood pressure ($\bar{x} \pm S D$) (mm Hg) | 179 \pm 29 | 182 \pm 22 | 184 \pm 21 |
| Diastolic blood pressure ($\bar{x} \pm S D$) (mm Hg) | 120 \pm 28 | 117 \pm 14 | 113 \pm 13 |
| Number with hypertensive retinopathy according to Keith-Wagener | | | |
| Stage I | 8 | 11 | 9 |
| Stage II | 9 | 5 | 7 |
| Stage III | 3 | 3 | 3 |
| Stage IV | 0 | 0 | 1 |
| Number with left ventricle hypertrophy on ECG | 9 | 7 | 8 |
| Number in WHO classification | | | |
| Stage I | 10 | 11 | 7 |
| Stage II | 5 | 3 | 6 |
| Stage III | 5 | 5 | 6 |
| Stage IV | 0 | 0 | 1 |

ceptives fasting in the morning recumbent, before rising after three to seven days admission to the hospital while on a daily sodium intake of 100 to 200 mEq. The urinary sodium excretion was measured during the 24 hours prior to this blood sampling and ranged in fact from

87 to 204 mEq with an average of 134 mEq. Ten ml blood were withdrawn in an ice-cooled tube containing 0.2 ml N,N -ethylenediaminetetraacetic acid (60 mg/ml solution) and centrifuged at 4°C. The plasma was deep frozen at -20°C. The renin concentration determined

attack are too small to substantiate or invalidate the Laragh hypothesis. Lacking a valid control series we classified our patients in three tertiles and compared the three groups.

Although our series is small, two patients with non malignant essential hypertension without heart failure and with low renin developed complications (stroke and heart attack). According to the suggestion of Brunner and colleagues¹ these patients should have been protected by their low renin.

III Comparison between the low renin groups of both series. In view of these divergent results we wonder what is the difference between the study of Brunner and colleagues¹ and the present study?

1. The selection was different. In our study only white patients and patients with essential hypertension were initially admitted and patients with heart failure or malignant hypertension were also included; however as discussed above exclusion of the latter patients would not basically influence the results.

2. Our patients with low plasma renin concentration had probably more severe hypertension; indeed while the diastolic blood pressure in our low renin group averaged 120 mm Hg, the diastolic blood pressure in Brunner and colleagues¹ low renin group averaged 105 mm Hg.

Furthermore the average creatinine clearance of the low renin group in our study was 79 ml per minute while in the study of Brunner and colleagues¹ it was 95 ml per minute.

The degree of retinopathy could not be compared since no exact figures were given in the paper of Brunner and colleagues¹. Left ventricular hypertrophy was present in 20 per cent of Brunner's and in 45 per cent of our low renin patients. Although the criteria for left ventricular hypertrophy may have been different, the greater percentage in our study is in accord with the higher diastolic pressure and lower creatinine clearance. The larger number of patients with more severe hypertension in our low renin group could explain the higher incidence of cardiovascular complications when compared with Brunner and colleagues¹ low renin group.

Based on Brunner and colleagues¹ communication one could speculate that a low renin level might protect against stroke and heart attack only if the hypertension was moderate but this of course remains to be confirmed.

3. Differences in methodology have also to be considered. Brunner and colleagues¹ measured plasma renin activity while the method used in our study (Skinner²) is thought to measure plasma renin concentration. Brunner and colleagues¹ drew blood at noon after their patients had been ambulatory for at least 4 hours while we drew blood in the morning, our patients having been recumbent overnight. Furthermore the mean urinary sodium excretion in Brunner and colleagues¹ low renin group was 16.1 mEq per 24 hours while the 24 hour urinary sodium excretion in our own study ranged between 87 and 204 mEq. It is not clear to us how these differences in methodology could explain the differences in cardiovascular complications.

4. The general approach in both studies was different. Brunner and colleagues¹ compared the plasma renin levels of their patients with those of normal volunteers while we classified our patients in three tertiles according to their plasma renin concentration.

We have already discussed the fact that Brunner and colleagues¹ control series did not contain any black subject but this would not invalidate their results if we consider only their patients and exclude their control group.

We then find in Brunner and colleagues¹ low renin group 27 per cent of their patients which is similar to 1/3 or 33 per cent of patients in our low renin group.

This difference in the classification system would not change our patients with stroke and heart attack from the low to the middle renin group since their plasma renin concentration was indeed very low as indicated in Fig. 1, namely 3 and 3.8 units per ml respectively.

IV General conclusion and prospectives. In conclusion in our study we did not find any data to confirm the hypothesis of Brunner and colleagues¹ concerning the protection against stroke and heart attack by a low plasma renin level. It is possible that

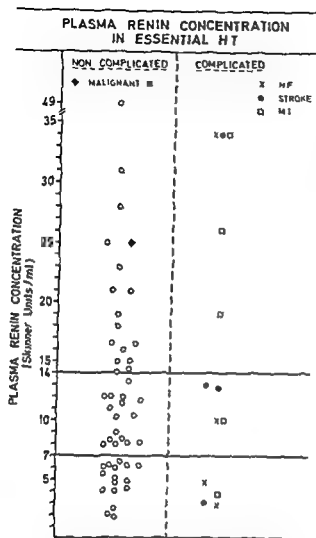


Fig. 1 Plasma renin concentration in essential hypertension. Each sign represents the plasma renin concentration of the respective 59 patients with essential hypertension. The patients represented in the right column under the heading, *COMPLICATED*, had at least one of the three following complications: heart failure (x), stroke (●) or myocardial infarction (□); the patient with two and the other with three complications at the same time are represented by two or three signs surrounded by a dotted line. The patients without any of these three complications are represented in the left column under the heading, *NON COMPLICATED*; they include patients with benign essential hypertension (○) except for one who had malignant hypertension (⊕). The two horizontal lines represent the dividing lines between the three groups of patients and not the limits of the normal range

in the low renin group we also retain in these circumstances two patients with stroke in the middle renin group and two patients with heart attack in the high renin group. In our study only one patient with malignant hypertension (defined by the presence of a high blood pressure and

papilledema) was entered as can be seen from Fig. 1 he had a high plasma renin concentration and no heart failure, heart attack or stroke. Withdrawing this one patient would therefore not influence the incidence of cardiac complications or stroke in any group.

Discussion

I Discussion of the paper of Brunner and colleagues¹ The study of Brunner and colleagues¹ performed in well standardized conditions of sodium intake in a large group of hypertensive patients with a big control series yielded provocative results and led to a discussion in the medical literature during the past year—in editorial in the *British Medical Journal*² the *Lancet*³ and a letter by Golby and associates⁷ and Schalekamp and co-workers.⁸ We will review these comments and add some more remarks.

A Brunner and colleagues¹ mentioned that there was no significant difference in the average diastolic blood pressure (DBP) between the group with low (DBP = 105 ± 14 mm Hg) and normal renin (DBP = 104 ± 17 mm Hg). They did not discuss however the variability of the blood pressure. Also the systolic blood pressure in itself could be considered as a risk factor.

B It is generally accepted that for the same blood pressure duration females have less cardiovascular complications than males. The authors did not mention in their paper the sex ratio in the three groups.⁹

C The percentage of black patients in the low renin group (42 per cent) was higher than in the normal renin group (24 per cent) furthermore the percentage of black subjects in the control group was not mentioned. Therefore we do not know if the control and the patient series are matched for this criterion.¹

II Discussion of the present paper The present study lacks the numbers and comparison with a control series when compared with the study of Brunner and colleagues.¹ Indeed in our study the numbers of patients with stroke or heart

According to a recent personal communication from Larashi the sex ratio was the same in the three groups.
The finding of a personal communication from Larashi the control group did not contain any black subject.

- 2 Skinner S L. Improved assay methods for renin concentration and activity in human plasma *Circ Res* 20:391 1967
- 3 Laragh J H. Biochemical profiling and the natural history of hypertensive diseases: low renin essential hypertension: a benign condition *Circulation* 44:971 1971
- 4 Laragh J H, Baer L, Brunner H R et al. Renin, angiotensin and aldosterone system in pathogenesis and management of hypertensive vascular disease *Am J Med* 52:633 1972
- 5 Editorial. New thoughts on essential hypertension *Br Med J* 2:121 1972
- 6 Editorial. Angiotensin, myocardial infarction and strokes *Lancet* 1:1273 1972
- 7 Golby F S and Berlin L J. New thoughts on essential hypertension *Br Med J* 2:594 1972
- 8 Schalekamp M A D H and Birkenhager W H. Renin levels in hypertension *N Engl J Med* 286:1320 1972

the higher complication rate in our low renin group could be related to the more severe hypertension in our group

A prospective study of this topic is needed to investigate several problems raised by these previous studies, but not yet clearly solved

(1) Should we consider these three groups as three distinct entities as suggested by Brunner and colleagues¹ or as three stages in the evolution of hypertension in the single patient as suggested by Schalekamp and associates² The impression of Brunner and colleagues¹ was that the patients did not change from group to group, although their follow up was too short to enable definite conclusions

(2) Furthermore the possibility should be excluded that a stroke or myocardial infarction may cause an increased plasma renin level the increase in renin therefore being the consequence rather than the cause of the cardiovascular complication

(3) Because both studies were retrospective the influence of previous treatment could not be evaluated It is indeed possible that previous antihypertensive treatment could have prevented cerebrovascular accidents in the low renin group although it is not clear if this treatment could also prevent heart attacks

(4) In an editorial Laragh³ suggested that antihypertensive agents which lower plasma renin levels (e.g. beta blocking agents methyl dopa clonidine, reserpine) should be preferred to those which increase the renin levels (e.g. diuretics) He did not present a direct proof of this statement It should be remembered however that although the influence of short term treatment of several antihypertensive agents on plasma renin levels is well known the influence on circulating renin of long term treatment with these drugs is often debated, because of conflicting results

Furthermore, diuretics which are known to increase plasma renin levels at least in acute experiments, have been used in combination with other antihypertensive agents in several studies which have shown a decreased incidence of strokes It is unlikely that the beneficial effect of this combined treatment was related exclusively to the non diuretic agents In a

more recent article Laragh and colleagues⁴ mentioned the possibility that the vascular toxicity supposedly due to increased renin levels, which could be suspected to be present during thiazide treatment, might be prevented by the diuretic action of these drugs themselves

As long as these problems have not been adequately solved in prospective studies therapeutic conclusions drawn from these data could be premature

Summary

Fifty nine patients with essential hypertension were divided into three tertiles based on their plasma renin concentration determined on blood withdrawn in standardized conditions of day physical activity sodium intake and drug treatment

The incidence of known risk factors for cardiovascular disease such as sex age family history cigarette smoking obesity systolic and diastolic blood pressure plasma cholesterol and lipids did not differ significantly between the 20 patients with low renin levels (below 7 Skinner units per milliliter) the 19 patients with middle range renin levels (between 7 and 14 U per milliliter) and the 20 patients with high plasma renin levels (14 units per milliliter or more)

The incidence of stroke and heart attack was the same in the three groups irrespective of whether patients with heart failure and with malignant hypertension were included or excluded from the series

We therefore could not find from our data a support for the hypothesis which suggests that low renin protects hypertensive patients against stroke and heart attack The differences between this study and the study of Brunner and associates¹ and Laragh and colleagues⁴ are discussed in an attempt to explain this divergent tendency It is felt that therapeutic conditions cannot be drawn from this small and retrospective study but that large scale prospective studies are needed

REFERENCES

- 1 Brunner H R Laragh J H Baer L et al Essential hypertension renin and aldosterone heart attack and stroke *N Engl J Med* 286 441 1972

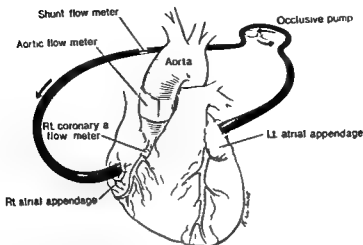


Fig 1 Diagram showing experimental preparation Right ventricular output increased by hunting blood from the left atrium to the right atrium through external conduit Left ventricular output and hunt flow were monitored with electromagnetic flow meters

Methods

We anesthetized 9 dogs weighing 16 to 28 kilograms with sodium pentobarbital 30 mg per kilogram of body weight intravenously and gave supplemental amounts when needed. The dogs were ventilated by a Harvard respirator through an endotracheal tube with oxygen enriched gas mixtures to give arterial oxygen tensions greater than 100 mm Hg and carbon dioxide tensions between 28 to 40 mm Hg. A bilateral thoracotomy was done through the fourth intercostal space. Catheters were placed in both femoral arteries, a femoral vein, the ascending aorta, left atrium, distal pulmonary artery, and directly into the right ventricle through the outflow tract. Pressures were measured with Statham P 23Db transducers and recorded by a Beckman oscillograph. A tourniquet of umbilical tape placed around the main pulmonary artery could be tightened to produce the desired level of right ventricular hypertension.

Number 34F Bardex tubes were inserted through the left and right atrial appendages (Fig 1). A 10 mm internal diameter (ID) cannulating electromagnetic flow transducer (Statham Model SP 2202) was placed in the tubing connecting the two atrial cannulae and the tubing was inserted into an occlusive roller pump to control the size of the interatrial left to right shunt. Aortic flow was recorded with 13 to 16 mm ID

electromagnetic flow transducers (Statham) placed around the ascending aorta and pulmonary flow was calculated as the sum of aortic and shunt flows. The flow transducers were calibrated *in vitro* with timed volume collections of blood. A 20 mm ID electromagnetic flow transducer (Statham) was placed around the proximal right coronary artery to measure changes in the post occlusive hyperemic response.⁴ Zero flow reference levels were determined by 8 sec and occlusions of the right coronary artery distal to the flow transducer. The flow debt during occlusion was calculated as baseline flow times the duration of occlusion. The payback of this debt was calculated by planimetry of the area beneath the mean coronary flow curve and above the baseline flow level from the time of release of the occlusion until mean flow stabilized at or near the pre occlusive level. The ratio of this payback area to the flow debt was multiplied by 100 to give the per cent hyperemic response. 100 per cent indicates that the flow debt was exactly repaid.⁴

We measured regional myocardial blood flow by injecting differently labeled batches of radioactive microspheres into the left atrium as described previously.^{5,6} Filtered microspheres with a mean diameter of 7 microns (range 4 to 12) labeled with ^{14}Ce , ^{86}Sr and ^{45}Sc were used to measure intra ventricular regional flow. However we also used 15 micron diameter microspheres

Regional coronary flow with increased right ventricular output in anesthetized dogs

David E Fivler MD
Joseph P Archie, Jr MD
Daniel J Ulliot, MD
Julien I E Hoffman MD
Dallas Texas and
San Francisco Calif

In many types of congenital heart disease intracardiac left to right shunts increase right ventricular output and frequently produce right ventricular dilatation and pulmonary hypertension. While the clinical effects of these shunts are well known their effect on right ventricular coronary flow has not been studied. Ventricular dilatation and elevated ventricular pressure produce an increase in wall tension hence a greater myocardial oxygen need. Whether these changes in myocardial oxygen need are accompanied by appropriate increases in right ventricular myocardial flow is not known.

Myocardial flow is determined by two main factors coronary vascular resistance and coronary driving pressure the latter for the right ventricle being defined as the difference between aortic and right ventricular pressures.¹ This means that with pulmonary hypertension, the increased right ventricular pressure tends to lower the

coronary driving pressure. As right ventricular coronary driving pressure falls coronary vasodilatation must occur to maintain right ventricular coronary flow. Recent studies of the right ventricle have indeed shown that the initial response to acutely increased right ventricular pressure is local vasodilatation.^{2,3} Following maximal coronary vasodilatation a further rise in right ventricular pressure lowers right ventricular coronary driving pressure and limits coronary flow. Therefore in left to right shunts right ventricular coronary flow would be adequate if there is coronary vasodilatation as right ventricular hypertension develops but would be inadequate if maximal coronary vasodilatation occurred while right ventricular coronary driving pressure was still decreasing. The purpose of this study was to determine the effects of acute short term increases in right ventricular output on coronary flow to the right ventricle.

From the Department of Pediatrics Southwestern Medical School University of Texas Dallas Texas, and the Cardiovascular Research Institute University of California San Francisco San Francisco Calif.
Supported in part by Program Project Grant HL 06285 from the National Heart and Lung Institute Clinical Investigators in Surgery T 01 GM 01924-03 (Dr Archie) National Institute of Health Special Fellowship HL 4342-02 (Dr Fivler) National Heart and Lung Institute Graduate Training Grant HL 5251 (Dr Ulliot) and the American Heart Association Texas Affiliate.
Received for publication March 5 1973.
Reprint requests to Dr David L. Fivler Department of Pediatrics Southwestern Medical School University of Texas 5323 Harry Hines Blvd Dallas Texas 75235.

December 1973

Table II Mean coronary flow values for the right and left ventricles

| | Control | Small shunt | Moderate shunt | Large shunt | Small shunt and RV hypertension |
|---|-------------|--------------|----------------|--------------|---------------------------------|
| REGIONAL MYOCARDIAL FLOW (ml/min. 100 Gm.) | | | | | |
| Right ventricular | | | | | |
| Free wall | 40.8 ± 15.8 | 8.8 ± 3.8† | 64.5 ± 21.0† | 94.2 ± 24.2† | 81.8 ± 33.4† |
| Septal | 9.2 ± 28.0 | 77.2 ± 36.5† | 64.0 ± 15.5† | 97.2 ± 28.4† | 94.3 ± 16.3† |
| Left ventricular | | | | | |
| Free wall | 64.0 ± 22 | 81.7 ± 32.3 | 58.5 ± 8.4 | 69.8 ± 22.0 | 69.3 ± 6.6 |
| Septal | 18.7 ± 23.3 | 84.0 ± 36.0 | 63.5 ± 12.1 | 84.5 ± 18.2 | 68.3 ± 28.0 |
| MYOCARDIAL FLOW DISTRIBUTION | | | | | |
| Right ventricular (Free wall) | | | | | |
| Subendocardial | | | | | |
| Subepicardial | 78 ± 04‡ | 77 ± 08 | 0 ± 16 | 53 ± 20 | 59 ± 11 |
| Left ventricular (Free wall) | | | | | |
| Subendocardial | | | | | |
| Subepicardial | 96 ± 10‡ | 90 ± 14 | 95 ± 29 | 58 ± 09 | 94 ± 21 |

Values are mean ± one standard deviation. Control values are averaged from all animals. Values are not averages of the same animals in each group. The effect of changes in flow are clear-cut in Fig. 1.

† $p < 0.05$ (Duncan's multiple range test).

‡ Values determined using 7 micro-diam. microsphere technique (Ref. 2).

measured by planimetry of the area beneath the systolic portion of the right ventricular pressure tracings. The left ventricular tension time index was measured by planimetry of the area beneath the systolic portion of the aortic pressure tracing as described by Sarnoff and colleagues.⁸ These indices are expressed as mm Hg sec/min that is in terms of total area per minute. All statistical comparisons were made by paired *t* testing and by analysis of variance.⁹

Results

The most prominent changes that occurred with large left to right shunts were an increase in right ventricular systolic and diastolic pressures, an increase in right ventricular output, and a fall in heart rate (Table I). The changes in right ventricular systolic pressure with pulmonary arterial constriction and with moderate and large left to right shunts were not statistically different from each other. Left ventricular output, aortic pressure, and left ventricular tension time index did not significantly

differ between altered states. Heart rate was significantly lower than control in all except the large shunt group, but did not differ significantly among the altered states.

Mean coronary flow values (ml/min. 100 Gm.) for the right and left ventricles are listed in Table II. Moderate variability in coronary flows is seen within each group even in the controls due to small differences in hemodynamics and oxygen contents. It is also important to point out that the flow values in Table II are not averages of the same animals in each of the five groups, since only four flow determinations could be made in any one animal. Therefore changes in coronary flows are illustrated more clearly in Fig. 2 where changes were analyzed with each animal serving as its own control (paired *t* testing). Coronary flow to the right ventricular free wall and to the right side of the septum increased initially ($p < 0.05$) with small and more with moderate left to right shunts. Coronary flow failed to increase progressively, however, with either large or small shunts having right ventricular hypertension (right ven-

Table I Findings at time of myocardial flow determinations

| | Control | Small shunts | Moderate shunts | Large shunts | Small shunts and RV hypertension |
|--|-------------|--------------|-----------------|--------------|----------------------------------|
| Number of determinations | 9 | 7 | 6 | 4 | 6 |
| Heart rate (beats/min) | 150 ± 9.3* | 121 ± 27.2† | 110.3 ± 28.1† | 127.5 ± 8.5 | 124 ± 31.7† |
| Right ventricular output (L/min) | 142 ± 0.36 | 307 ± 0.70† | 387 ± 0.80† | 572 ± 0.98† | 338 ± 0.6† |
| Left ventricular output (L/min) | 142 ± 0.36 | 160 ± 0.46 | 162 ± 0.46 | 210 ± 0.76† | 155 ± 0.35 |
| Right ventricular systolic pressure (mm Hg) | 70 ± 6.87 | 377 ± 2.67† | 377 ± 9.32† | 450 ± 7.55† | 547 ± 9.6† |
| Right ventricular diastolic pressure (mm Hg) | 67 ± 1.83 | 100 ± 0.93† | 107 ± 3.14† | 113 ± 1.45† | 137 ± 3.94† |
| Right ventricular tension time index (mm Hg/min) | 606 ± 175 | 980 ± 174† | 1050 ± 279† | 1030 ± 353† | 1550 ± 341† |
| Aortic mean pressure (mm Hg) | 85.2 ± 11.0 | 76.7 ± 26.1 | 78.8 ± 11.0 | 88.8 ± 15.9 | 70.0 ± 20.8 |

*Values are means ± one standard deviation

†P < 0.05 (Significant difference from control)

labeled with ^{125}I to measure interventricular myocardial flows. A small Holter pump was used to withdraw reference blood samples (timed volume collections) from the femoral arteries. Since well mixed microspheres are distributed to organs or regions in proportion to regional flow, regional myocardial flow was determined from the equation: myocardial flow/myocardial nuclide activity equals reference sample flow/reference sample nuclide activity. At the end of each experiment the dog was killed and the heart was removed. The atria, valves, great arteries, large coronary vessels, and epicardial fat were removed from the ventricles. The free walls of the right and left ventricles were cut from the septum and divided into 3 layers (subendocardial, middle, and subepicardial regions) of about equal thickness. The ventricular septum was divided into three layers: right, middle, and left. The tissue from each region was placed in one or more vials, weighed, and counted in a well scintillation counter. The radioactivity emitted from each nuclide was determined by the method of Rudolph and Heymann, modified by using the reference sample technique to calculate flows and by changing the constants used for differential spectrometry after recalibrating the counting equipment.⁷

Regional myocardial flow was measured during control state (no left to right shunt)

in each dog with 15 micron microspheres. Pulmonary flow was then increased by 100 per cent, 200 per cent, and 300 per cent (small, moderate, and large shunts) in random sequence. In 5 dogs pulmonary flow was increased by 150 per cent and right ventricular hypertension (peak right ventricular pressure greater than 45 mm Hg) was produced by pulmonary arterial constriction. With all shunts Ringer's lactate and/or fresh blood (125 to 275 ml) were infused to maintain left ventricular output and aortic pressure at control levels. Changes in myocardial flow were determined with each dog serving as its own control.

Total coronary vascular resistance was calculated from the equation: resistance equals pressure difference/flow, where the pressure difference was the coronary driving pressure, and flow was the myocardial flow (per 100 grams) as measured with radioactive microspheres. This resistance calculation includes the effects of increased extravascular compression during systole as well as a coronary vasomotor tone. The coronary driving pressure index (PI) of the right ventricle was calculated by planimetry of the area (per minute) between the aortic pressure tracing and the right ventricular pressure tracing. Oxygen demand of the right ventricular myocardium was estimated from its tension time index (TTI)

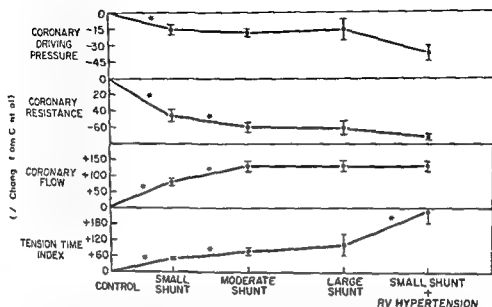


Fig 3 Per cent changes in right ventricular coronary hemodynamics in the four shunt groups. Per cent changes are calculated from each dog's own control value. Each point represents mean per cent change. Vertical brackets indicate the standard error of the mean. Asterisks indicate significant difference ($p < 0.05$) between adjacent points.

124 per cent \pm 13. This indicates that when right ventricular output and right ventricular pressure increased, coronary vasomotor reserve was diminished.

Changes in right ventricular coronary hemodynamics are shown in Fig 3. Coronary driving pressure progressively fell as right ventricular hypertension developed: it averaged 4.37 ± 30.8 (SE) mm Hg per minute for the control group and 3.822 ± 366 mm Hg per minute, 3.420 ± 276 mm Hg per minute, 3.360 ± 570 mm Hg per minute and 2.700 ± 440 mm Hg per minute in the 4 shunt groups respectively. As coronary driving pressure fell, coronary vasodilation occurred as reflected by the fall in coronary vascular resistance. However, coronary resistance did not fall significantly further in the groups with large or small shunts having right ventricular hypertension and this suggests that the coronary arteries were maximally vasodilated in these groups. Right ventricular coronary flow increased as coronary resistance fell, but in groups with right ventricular hypertension failed to increase further when coronary vascular resistance failed to change. Right ventricular tension time index was increased by the elevation

in right ventricular systolic pressure and the prolongation of systole with greater stroke volumes. With shunts having only mild elevation in right ventricular pressure (small and moderate groups), coronary flow increased as the tension time index increased. In shunts with right ventricular hypertension, myocardial flow failed to increase further as the tension time index rose.

To determine if we could predict from the pressure tracings when the right ventricular coronary arteries were no longer able to increase flow, we related the right coronary driving pressure index (PI) to the right ventricular tension time index (TTI), that is estimated oxygen supply potential compared to estimated oxygen needs.² This PI/TTI ratio averaged 7.5 ± 0.72 (SE) in the control group and fell significantly ($p < 0.05$) with small left to right shunts but thereafter did not significantly change, averaging 3.5 ± 0.60 , 3.7 ± 0.46 and 3.4 ± 0.88 respectively. In the group with small shunts and right ventricular hypertension, the PI/TTI ratio fell to 1.7 ± 0.29 , the difference from the other shunt groups being just above the conventional level of significance ($p < 0.10$).

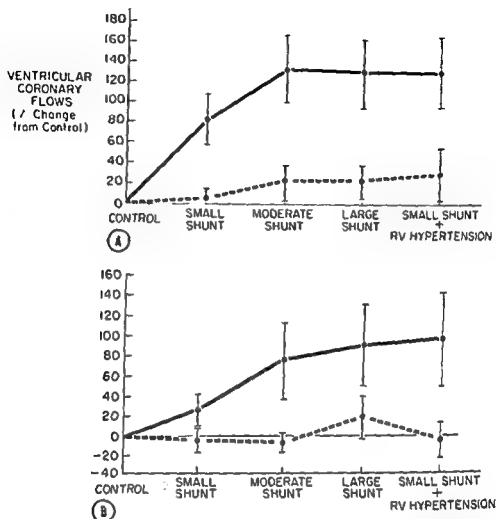


Fig 2 A and B Per cent changes in myocardial blood flow per gram to the free wall (A) and septal region (B) of the right (solid line) and left (dashed line) ventricles in the four shunt group. Per cent change is calculated from each dog's own control value. Small, moderate and large shunts had pulmonary flow increased by 100 to 150 per cent, 200 per cent and 300 per cent respectively. RV hypertension is defined as right ventricular peak pressure above 45 mm Hg. Each point represents mean per cent change; vertical brackets the standard error of the mean.

tricular peak pressure greater than 45 mm Hg). Coronary flow to the left ventricular free wall and the left side of the septum did not change significantly from control. Changes in flow to the right and left portions of the ventricular septum resembled those to their respective free walls (Fig. 2). The regional distribution of myocardial flow within the free walls of both ventricles as reflected by the ratio of subendocardial flow:subepicardial flow (per 100 Gm) did not change significantly (Table II). The distribution of coronary flow between the ventricles did change: for the control group myocardial flow (per 100 Gm) to the free wall of the right ventricle was 60 per cent of that in left ventricle. In all of the shunt

groups myocardial flow (per 100 Gm) to the free wall and septal regions of the right ventricle equalled or exceeded that to the left ventricle. This change in coronary flow distribution was due to the increase in right ventricular coronary flow since left ventricular coronary flow remained unchanged.

To estimate changes in coronary vaso motor reserve we measured the reactive hyperemic response of the right coronary artery at different right ventricular outputs. In controls the occlusive coronary flow debt was repaid by 150 per cent \pm 9.7 (SE), whereas with progressively larger shunts it averaged 121 per cent \pm 10, 110 per cent \pm 12, 127 per cent \pm 14, and with small shunts and right ventricular hypertension

Coronary flow to the septal portion of the right ventricle increased significantly with greater right ventricular volume loads. With effective autoregulation this indicates that septal myocardial oxygen consumption increased and suggests that septal contraction plays an important role in maintaining right heart output. Other studies have suggested that the ventricular septum is responsible for maintenance of right ventricular performance following functional removal of the right ventricular free wall.^{11,12} Most recently Brooks and co-workers¹³ reported that complete occlusion of the right coronary artery in dogs caused no change in cardiac output or right ventricular pressure but that with the added stress of elevated right ventricular pressure right heart failure occurred. In the present study once right ventricular pressure became elevated septal coronary flow failed to increase further. This suggests that right ventricular hypertension may cause myocardial compression of coronary vessels within the ventricular septum thereby impeding coronary flow to this region.

One purpose of this study was to determine if acute right ventricular volume loads exhaust right ventricular coronary reserve. With small left to right shunts a slight increase in right ventricular pressure reduced coronary driving pressure however coronary flow was increased by vasodilatation as indicated by the fall in coronary vascular resistance (Fig. 3). In fact the maximal increase in right ventricular myocardial flow occurred after right ventricular output was increased by 200 per cent (Fig. 2). With larger left to right shunts or when right ventricular systolic hypertension developed myocardial flow failed to increase further. Under these circumstances coronary vascular resistance failed to fall further suggesting that the coronary vessels were fully dilated and that this compensatory mechanism could no longer increase coronary flow. The decreased reactive hyperelemic responses of the right coronary artery also indicated that coronary vasomotor reserve was reduced. In previous studies of acute right ventricular hypertension alone we found that myocardial flow initially increased with relatively mild

hypertension but failed to increase further with moderate hypertension.² Aukland and co-workers³ also reported that right ventricular hypertension alone increased right ventricular myocardial flow but usually less 200 per cent of control. This may be explained on the basis that right ventricular hypertension decreases coronary driving pressure and limits coronary flow. We found that acute cardiovascular collapse characterized by precipitous fall in blood pressure and systemic flow frequently occurred at those levels of right ventricular hypertension at which myocardial flow failed to increase. Under such conditions the greater myocardial oxygen needs were not met by increased oxygen supply and resulted in relative coronary insufficiency. In the present study myocardial flow failed to increase further with greater pressure and volume loads suggesting that an acute left to right shunt with right ventricular hypertension may also result in relative coronary insufficiency.

It would be most helpful if acute cardiovascular collapse secondary to right ventricular hypertension could be predicted from hemodynamic data. In our study of acute right ventricular hypertension alone we found the relationship between the right coronary artery driving pressure index and its tension time index was useful in predicting when right coronary vessels were unable to increase flow.² This PI/TTI ratio compared the pressure aspect of myocardial oxygen supply to the pressure aspect of myocardial oxygen need. The relationship between the tension time index and myocardial oxygen uptake however is affected by changes in inotropic state, heart rate and ventricular geometry and volume. In the present study right ventricular diastolic pressure increased significantly with the acute changes in right ventricular output and indicated that right ventricular volumes probably increased. Under such circumstances the tension time index may not reflect the effect of larger right ventricular volume on true wall tension and hence probably underestimates changes in myocardial oxygen need. In the acute right ventricular hypertension study right ventricular myocardial flow failed to increase when the PI/TTI ratio averaged 2.4 ± 0.27 .

Discussion

To our knowledge this is the first report describing the effects of acute right ventricular volume and pressure overloading on regional coronary flow. The present study provides evidence that coronary flow to the right ventricle is predominantly influenced by factors which alter pressure work rather than flow work and suggests that wall tension in the left ventricle is probably one of the main determinants of oxygen need and thus regional coronary flow. It is important to consider how tension in the right ventricular wall changes with different pressure and volume loads.

Theoretically according to the Laplace equation wall tension is directly proportional to both cavity pressure and radius of curvature. Assuming that the right ventricular free wall is nearly hemispherical then if peak pressure is doubled at constant radius wall tension doubles. On the other hand if stroke volume is doubled at constant peak pressure only a small increase occurs in the radius of curvature and therefore there should be only a small increase in wall tension. This is what was found in the left ventricle by Braunwald and associates¹⁰ changes in left ventricular peak pressure resulted in greater alterations of myocardial oxygen consumption than did changes in cardiac output. When cardiac output was increased by 100 per cent and 200 per cent at constant peak left ventricular systolic pressure and heart rate coronary flow increased approximately 10 per cent and 20 per cent respectively (calculated from their Fig. 5). This is consistent with the study of Taylor and colleagues¹¹ who found that when left ventricular stroke volume doubled at constant peak pressure peak wall tension increased by only 18 per cent.

In the present study when right ventricular output was increased by 100 per cent and 200 per cent myocardial flow increased by 81 per cent and 134 per cent (Fig. 2). Right ventricular systolic pressure concurrently increased from 27 mm Hg to 38 mm Hg indicating that right ventricular myocardial flow might have increased from the effects of greater pressure as well as from the effects of the larger ventricular output. Unfortunately in the present study

we could not determine the extent to which right ventricular volume overload alone increased right ventricular myocardial flow. However, in recent studies of acute right ventricular hypertension we found that when right ventricular systolic pressure increased in a similar fashion to the present study (from 26 mm Hg to 37 mm Hg) right ventricular myocardial flow (per 100 grams) increased from 65.7 ± 6.2 to 79 ± 7.2 ml per minute (mean and standard error).² The latter flow value is not significantly different from the average right ventricular myocardial flow values obtained in groups with small and moderate shunts (75.8 and 65.4 ml/min/100 gm). These data suggest that like the left ventricle the increase in ventricular pressure was principally responsible for the increase in right ventricular coronary blood flow.

This study demonstrates that an increased volume load to the right ventricle resulted in an increase in right ventricular myocardial flow while left ventricular myocardial flow remained unchanged (Fig. 2). Earlier studies have shown that coronary flow to the right and left ventricles may vary independently each being determined mainly by the pressure load of the respective ventricle.^{2,3} Changes in the inter-ventricular distribution of coronary blood flow therefore seem to reflect the work loads of the respective ventricles rather than the anatomical distribution of the coronary arteries. In dogs the left coronary artery supplies approximately one third of the right ventricular free wall.¹² As the pressure and volume load of the right ventricle increased a greater proportion of total coronary flow went to the right ventricle presumably through both the right and left coronary arteries. If the changes in regional coronary flow resulted from altered sympathetic tone then an increase in left coronary flow might be expected to increase myocardial flow in both the left and right ventricles. However left ventricular myocardial flow did not change significantly indicating that coronary vascular resistance maintained left ventricular flow in accordance with its metabolic requirements. This suggests that autoregulation of coronary flow is regional, being determined by regional metabolic factors.

- consumption and coronary flow *Am J Physiol* 192:157 1958
- 11 Taylor R R, Cingolani H F and McDonald R H Jr Relationships between left ventricular volume ejected fraction and wall stress *Am J Physiol* 211:674 1966
- 12 Blair E Anatomy of the ventricular coronary arteries in the dog *Circulation* 9:333 1961
- 13 Starr I, Jeffers W A and Mead R H The absence of conspicuous increments in venous pressure after severe damage to the right ventricle of the dog with a discussion of the relation between clinical congestive failure and heart disease *Am Heart J* 26:291 1943
- 14 Bakes A C P The question of the function of the right ventricular myocardium: an experimental study *Circulation* 11:724 1950
- 15 Brooks H I, Kirk E S, Vokonas P S, Urthel C W and Sonnenblick E H Performance of the right ventricle under stress: relation to right coronary flow *J Clin Invest* 50:2176 1971
- 16 Sandier H and Dodge H T Left ventricular tension and stress in man *Circ Res* 13:91 1963
- 17 Frank M J, Nadimi M, Moschos C B and Levinson G E Left ventricular coronary flow, metabolism and performance in mild congenital heart disease with increased left ventricular flow or pressure *Am Heart J* 79:20 1970

(SE) whereas in the present study myocardial flow failed to increase further when this ratio averaged 3.7 ± 0.65 (moderate shunt group). Since the denominator of this ratio underestimates myocardial oxygen needs in this acute shunt study it seems reasonable to expect the ratio to be higher at the point where coronary flow is limited. Although less sensitive in the case of acute ventricular volume changes the ratio of PI/TI may provide a means for indirectly assessing right ventricular coronary reserve in the presence of acute left to right shunts.

Our results obtained with acute right ventricular volume overload in open chest anesthetized dogs cannot be applied directly to patients with chronic right ventricular overload and ventricular hypertrophy. Not only will there be differences in autonomic and humoral influences but in the hypertrophied ventricle the thicker wall reduces wall tension for any ventricular pressure and radius.¹⁶ Furthermore in congenital lesions there could be more myocardial vascularity and thus differences in pressure flow relations in the regional coronary bed. The importance of these differences between acute and chronic right ventricular volume overloads has not been evaluated but in one study of patients with left ventricular volume overloads Frank and colleagues¹⁷ reported that an increase in myocardial oxygen consumption occurred only in association with increases in the left ventricular tension time index. They suggested that in shunt lesions the stimulus which results in increased metabolic demand of the myocardium is best described as a pressure overload. If this is true also of right ventricular volume overloads, then the principles found in the present study even if not the numerical results, may well apply in human disease.

Summary

In this study of acute right ventricular volume overloading in anesthetized dogs right ventricular coronary blood flow increased probably in response to the effect of increased right ventricular pressure on myocardial oxygen needs. Left to right shunts with small elevations in right ventricular pressure had compensatory coro-

nary vasodilatation which increased coronary flow. However shunts with right ventricular hypertension had no further fall in coronary resistance and failed to have an additional increase in coronary flow. This suggests that with acute right ventricular volume overloading the presence of right ventricular hypertension increases myocardial oxygen needs but limits the ability of the coronary vessels to increase flow. Should myocardial oxygen needs increase and the coronary vessels be unable to increase myocardial perfusion relative coronary insufficiency may occur.

We gratefully acknowledge the technical assistance of Elizabeth Shipkin and Kap Saunderson.

REFERENCES

1. Cross C E. Right ventricular pressure and coronary flow. *Am J Physiol* 202:12 1962.
2. Fixler D E, Archie J P, Ulfhot D J, Buckberg G D and Hoffman J I. Effects of acute right ventricular systolic hypertension on regional myocardial blood flow in anesthetized dogs. *Am Heart J* 85:491 1973.
3. Aukland K, Kul F and Kjekshus J. Relationship between ventricular pressures and right and left myocardial blood flows. *Acta Physiol Scand* 70:116 1967.
4. Colfman J D and Gregg D E. Reactive hyperemic characteristics of the myocardium. *Am J Physiol* 199:1143 1960.
5. Domenech R J, Hoffman J I, Noble M I, Saunders K M, Henson J R and Subijanto S. Total and regional coronary blood flow measured by radioactive microspheres in conscious and anesthetized dogs. *Circ Res* 25:581 1969.
6. Buckberg, G D, Luck J C, Payne B D, Hoffman J I, Archie J P and Fixler D E. Some sources of error in measuring regional blood flow with radioactive microspheres. *J Appl Physiol* 31:598 1971.
7. Rudolph A M and Heymann M A. Circulation of the fetus in utero: methods for studying distribution of blood flow, cardiac output and organ blood flow. *Circ Res* 21:163 1967.
8. Sarnoff S J, Braunwald E, Welch G H Jr, Case R M, Stansby W N and Macruz K. Hemodynamic determinants of oxygen consumption of the heart with special reference to the tension time index. *Am J Physiol* 192:118 1958.
9. Snedecor G W and Cochran W G. Statistical methods. Ames 1967. The Iowa State University Press.
10. Braunwald E, Sarnoff S J, Case R M, Stansby W N and Welch G H Jr. Hemodynamic determinants of coronary flow: effect of changes in aortic pressure and cardiac output on the relationship between myocardial oxygen

- consumption and coronary flow *Am J Physiol* 192 157 1958
- 11 Taylor R R Cingolani H E and McDonald R H Jr Relations between left ventricular volume ejected fraction and wall stress *Am J Physiol* 211 674 1966
- 12 Blair E Anatomy of the ventricular coronary arteries in the dog *Circulation* 9 333 1961
- 13 Starr I Jeffer W A and Mead R H The absence of conspicuous increments in venous pressure after severe damage to the right ventricle of the dog with a discussion of the relation between clinical congestive failure and heart disease *Am HEART J* 26 291 1943
- 14 Bakos A C P The question of the function of the right ventricular myocardium: an experimental study *Circulation* 1:724 1950
- 15 Brooks H L Kirk E S Vokonas P S Urschel C W and Sonnenblick E H Performance of the right ventricle under stress: relation to right coronary flow *J Clin Invest* 50 2176 1971
- 16 Sandler H and Dodge H T Left ventricular tension and stress in man *Circ Res* 13 91 1963
- 17 Frank M J Nadimi M Moschos C B and Levinson G E Left ventricular coronary flow metabolism and performance in mild congenital heart disease with increased left ventricular flow or pressure *Am HEART J* 79 20 1970

Experimental myocardial infarction in the closed-chest dog Controlled production of large or small areas of necrosis

Michael V Cohen, M D *

Per Eldh, M D **

Boston, Mass

Because it has been impractical to study many aspects of coronary artery disease in man investigators have used animal models particularly the dog. Open chest methods of obstructing coronary arteries have been employed most frequently. Early attempts involved simple ligation of coronary arteries¹ or injection of a starch suspension into the aortic root. Because of the high mortality rate attendant upon sudden occlusion of coronary arteries methods of gradual yet progressive obstruction of vessels were developed. Ameroid constrictors,^{2,3} inflatable balloons,⁴ adjustable hydraulic occlusive devices,⁵ microspheres injected via an indwelling catheter in the left coronary artery,⁶ sutures and anastomoses around the coronary vessels,^{7,8} and thrombogenic wires introduced through the wall of the coronary artery⁹ have been employed to this end. The need for thoracotomy is a major drawback of these techniques and prompted the development of several closed chest methods.

The availability of selective angiographic

methods has greatly simplified the introduction of catheters into the coronary arteries. Selective embolization of the coronary artery with wire coils¹⁰ or steel cylinders,¹¹ steel balls,¹² lead foil,¹³ radiopaque spheres,^{14,15} microspheres,^{16,17} lyophilized spores,¹⁸ catheter tips,¹⁹ plastic cones,²⁰ thrombus²¹ or mercury^{22,23} has been accomplished. Furthermore ischemia has been induced by wedging catheters into distal coronary arteries^{24,25} and inducing thrombus formation with current passing from an electrode introduced selectively into a coronary artery.^{26,27}

We have modified one of these methods—that of Hammer and Piskunov²⁸ to allow either immediate or delayed occlusion of a coronary artery to permit assessment of the effect on mortality rate and infarct size.

Method

Adult mongrel dogs are anesthetized with sodium pentobarbital (30 mg per kilogram of body weight). After obtaining a control coronary angiogram with Reno

From the Cardiovascular Division, Department of Medicine and the Department of Radiology, Harvard Medical School and the Peter Bent Brigham Hospital, Boston, Mass.

Supported by United States Public Health Service Grants HL 05679, HL 16608 and GM 18674.

Received for publication March 12, 1973.

Reprint requests to Dr. Michael V. Cohen, Peter Bent Brigham Hospital, 721 Huntington Ave., Boston, Mass. 02115.

*Research Fellow in Medicine, Harvard Medical School and Peter Bent Brigham Hospital, Boston, Mass.

**Visiting Assistant Professor of Radiology, Harvard Medical School and Visiting Radiologist, Peter Bent Brigham Hospital, Boston, Mass.

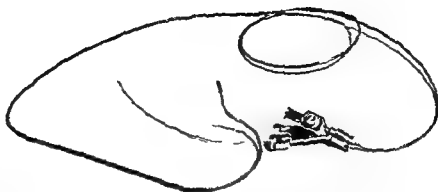


Fig. 2 Coronal catheter for embolization of coronary arteries with solid plugs. A guide wire has replaced the inner Teflon catheter.

Table 1 Survival related to type of plug

| Plug | Vessels | | |
|----------|---------|-----|--------------|
| | LAD* | LCf | First septal |
| Hollow | | | |
| Survived | 22 | — | 2 |
| Died | 3 | — | — |
| Solid | | | |
| Survived | 8 | 2 | 1 |
| Died | 2 | — | — |

* Abbreviations: LAD = left anterior descending artery
LCf = left circumflex artery

Table II Infarct size related to type of plug

| Dog | Vessel | Plug | % infarct |
|---------|--------|--------|-----------|
| 242 | LAD* | hollow | 0 |
| 338 | LAD | hollow | 16 |
| 342 | LAD | hollow | 0 |
| 344 | LAD | hollow | 0 |
| 419 | LAD | hollow | 0 |
| 475 | LAD | hollow | 5 |
| 520 | LAD | hollow | 6 |
| 569 | LAD | hollow | 5 |
| Average | | | 4 ± 2 |
| 48 | LAD | solid | 14 |
| 162 | LAD | solid | 24 |
| 176 | LAD | solid | 28 |
| 184 | LAD | solid | 18 |
| 339 | LCf | solid | 17 |
| 748 | LAD | solid | 21 |
| Average | | | 20 ± 2 |

* Abbreviations: LAD = left anterior descending artery
LCf = left circumflex artery

angiography. After acute interventions (to be reported elsewhere¹⁴) the animals were killed and the hearts were excised. The ventricular cavities were filled with gauze sponges to prevent collapse of the ventricles and the hearts were frozen. The hearts were sectioned every 4 to 5 mm from apex to base. The slices were photographed and the areas of visible infarction and total left ventricular surface were measured from the photographs by planimetry. It was quite simple to outline the necrotic areas which had a different texture and color compared to the surrounding myocardium. The ratio of infarct size to total left ventricular area was calculated.

Results

Hollow plugs were embolized into the coronary arteries of 27 dogs and solid plugs were used in 13 animals (Table I). There was an 11 per cent mortality rate associated with the hollow plugs: two dogs succumbed within one hour and one died after 48 hours. Two dogs with occlusive plugs died within one hour. There were no late deaths. All animals except one had well developed collateral vessels by the time of the repeat coronary angiogram (Fig 3). Twenty three of the surviving 24 dogs with hollow plug emboli had angiographic documentation of complete occlusion of the embolized vessel. The remaining dog had a patent LAD after three weeks despite apparent normal clotting mechanisms. A second hollow plug in the LAD resulted in complete occlusion.

The hearts of 14 dogs were serially sectioned, eight with hollow and six with solid



Fig 3 A and B Selected films from coronary angiograms done before (A) and three weeks after embolization (B) of the left anterior descending (LAD) coronary artery with a hollow plug. The LAD is totally occluded at the level of the plug and the distal LAD is filled retrogradely by left-to-left collaterals from the left circumflex and diagonal branches.

plugs (Table II). Seven of the eight animals with delayed coronary artery occlusion had either no or minimal visible areas of infarction (≤ 6 per cent). The eighth dog had 16 per cent of the left ventricle infarcted. Of animals with solid plugs all had transmural damage and an area of infarct ranging from 14 to 28 per cent (average 20 ± 2 per cent) of the measured left ventricular cross sectional surface (Fig 4). The difference in infarct size in dogs embolized with hollow plugs compared to those embolized with solid plugs is highly significant ($p < 0.001$).

After embolization of a hollow plug in two dogs coronary angiography was performed every 15 to 30 minutes until the plug occluded. Occlusion occurred between the fourth and fifth hours after embolization.

Discussion

Although animal models of coronary artery disease are not ideal many investigators have occluded coronary arteries in animals in an attempt to simulate some of the sequelae of naturally occurring coronary artery obstruction. Open-chest tech-

niques require thoracotomy with its added mortality rate and inconvenience. Adhesions between pericardium, anterior chest wall and myocardium make subsequent reoperation for study of collaterals or myocardial function more difficult and occasionally impossible. Furthermore the interruption of neural structures during coronary dissection may affect subsequent physiologic responses. Most closed chest methods require equipment not readily available in most radiology and catheterization laboratories: wire coils,¹² steel cylinders,^{14,15} steel balls,¹⁶ lead foil,¹⁷ radiopaque spheres,^{18,19} or microspheres.^{20,22} The technique described above however uses only Kifa and Teflon catheters and guide wires which are readily available. The technique is sufficiently versatile to allow for immediate or gradual occlusion with only minor alterations in approach. No method previously described permits this choice.

All described fluoroscopic methods save one²² have used the carotid artery. We have chosen the femoral approach. Selective coronary catheterization is much easier using this route and our preformed cath-



Fig 4 Serial cross sections of the left ventricle of a dog three weeks after embolization of the left anterior descending coronary artery with a solid plug. The macroscopic areas of necrosis are easily outlined. This dog had 24 per cent of its left ventricular cross sectional area affected by the abrupt deprivation of arterial blood.

eters. The femoral approach also allows the investigator to sacrifice both femoral arteries in the course of multiple angiographic, radioisotopic or other studies and use the carotid vessels in subsequent acute experiments. However, if reflex control and neurologic function are to be undisturbed, only one carotid artery can be sacrificed safely, thus limiting the number of possible interval studies. This is an important consideration in protocols employing multiple study points.

There has been meager documentation of the extent of infarction by any of these methods. Kordenat and associates¹² estimate 25 per cent infarction with wire coils lodged in the proximal LAD or LCx and 10 per cent infarction with coils in the distal vessels. Dogs with an inflatable balloon encircling the LAD just past its origin had infarction of 22 to 49 per cent of the left ventricle.⁸ Schaper²⁷ has noted that immediate LAD occlusion produces infarcts ranging from 11 to 33 per cent of the left ventricular mass, depending on the measurement technique employed, whereas approximately 50 per cent of dogs with gradual coronary occlusion from Ameroid constrictors have no infarcts. Ninety per cent (7/8) of our dogs with hollow plugs

had essentially no visible infarction of the left ventricle (average 2 per cent). Because of resorption of necrotic muscle, these numbers likely underestimate the size of the infarcted area. Four weeks after LAD occlusion, the thickness of the left ventricular wall in the area of the infarct may be decreased by up to 50 per cent.²⁷

Serial coronary angiography suggests that the hollow plug is occluded by thrombus at approximately four hours. The occlusion is unaccompanied by arrhythmias and often there is no characteristic change in the surface electrocardiogram. This gradually progressive coronary artery stenosis creates an area of ischemic myocardium which apparently induces the enlargement of already existing collateral pathways.^{28,29} Although these existing collaterals are not sufficient to supply the total metabolic requirements of the involved area, most probably they do serve a protective function and prevent significant areas of myocardium from becoming necrotic after complete occlusion of the embolized vessel. This hypothesis is supported by the observation of Elliot and colleagues²⁹ who gradually narrowed the LCx of dogs over 3 to 4 day periods and observed progressive increases of collateral indices during the con-

striction period and no or minimal evidence of infarction when the animals were killed. Thus this preparation is ideal for the study of collateral vessels or viable potentially ischemic myocardium. One surprising observation is the clear-cut protection afforded the dog by a relatively short period of gradually progressive stenosis before total coronary artery occlusion. Further studies are required to define the minimal duration of stenosis necessary and the exact nature of this protection in biochemical and hemodynamic terms.

The eighth animal with a hollow plug had 16 per cent of its left ventricular cross sectional surface area infarcted and it is postulated that the hollow plug thrombosed shortly after its embolization. Those dogs with solid plugs in the LAD averaged 20 per cent infarction. These animals were unable to profit from the enlargement of existing anatomic but non functional collateral pathways before the myocardium was irreversibly damaged.^{40, 41}

This model is suitable for examining post infarction hemodynamics, contractility, therapeutic modalities, etc. It is reasonable to suppose that larger infarcts and even shock models might be created by more proximal coronary artery obstruction with wider Kifa catheter plugs or with multiple plugs.

Summary

In order to embolize the coronary arteries a closed-chest technique was employed using fluoroscopically positioned catheters introduced selectively into the coronary arteries. Plugs made of catheter material were extruded directly into the coronary lumen. Small alterations in the coaxial catheter system permitted the use of either hollow or solid plugs with production of delayed or immediate coronary artery occlusions respectively. With hollow plugs the mortality rate was 11 per cent. An average of 4 ± 2 per cent of the left ventricular cross sectional area was infarcted. Solid plugs were associated with a mortality rate of 15 per cent and infarction of 20 ± 2 per cent of the left ventricle. Using this method it is therefore possible to create reproducible controlled infarct sizes in dogs enabling a careful study of collaterals

ischemic myocardium and therapeutic interventions

We would like to thank Ms. Patricia M. Sullivan R.T. and Ms. Catherine Favreau R.T. for their invaluable assistance in the angiographic studies and Drs. Edward S. Kirk and Herbert L. Abrams for their many helpful suggestions and criticisms.

REFERENCES

- Harris A.S. Delayed development of ventricular ectopic rhythms following experimental coronary occlusion. *Circulation* 1:1318 1950
- Roos A. and Smith J. III. Production of experimental heart failure in dogs with intact circulation. *Am J Physiol* 153:558 1948
- Litvak J., Sidendes L.E. and Vineberg A.M. The experimental production of coronary artery insufficiency and occlusion. *AM HEART J* 53:505 1957
- Vineberg A., Mahanti B. and Litvak J. Experimental gradual coronary artery constriction by ameroid constrictors. *Surgery* 47:765 1960
- Hood W.B. Jr., Jonson J., Kumar R., Katayama I., Neuman R.S. and Norman J.C. Experimental myocardial infarction I. Production of left ventricular failure by gradual coronary occlusion in intact conscious dogs. *Cardiovasc Res* 4:773 1970
- Khouri E.M., Gregg D.E. and Lowensohn H.S. Flow in the major branches of the left coronary artery during experimental coronary insufficiency in the unanesthetized dog. *Circ Res* 23:99 1968
- Weber K.T., Mahan T.I., Dennison B.H., Fuqua J.M. Jr., Speaker D.M. and Hastings F.W. Experimental myocardial ischemia and infarction. Production of diffuse myocardial lesions in unanesthetized calves. *Am J Cardiol* 29:793 1972
- Allen J.B. and Laadt J.R. The effect of the level of the ligature on mortality following ligation of the circumflex coronary artery in the dog. *AM HEART J* 39:273 1950
- Rushmer R.F., Watson N., Harding D. and Baker D. Effects of acute coronary occlusion on performance of right and left ventricles in intact unanesthetized dogs. *AM HEART J* 66:522 1963
- Hill J. III., Malinow M.R., McNulty W.P. and Ochsnor A.J. III. Experimental myocardial infarction in unanesthetized monkeys. *AM HEART J* 84:82 1972
- Mendlowitz M., Schauer G. and Gross L. Hemodynamic studies in experimental coronary occlusion II. Closed chest experiments. *AM HEART J* 11:664 1937
- Blair E., Nygren E. and Cowley H.A. A spiral wire technique for producing gradually occlusive coronary thrombosis. *J Thorac Cardiovasc Surg* 48:476 1964
- Kordenat R.K., Kezdi P. and Stanley E.L. A new catheter technique for producing experimental coronary thrombosis and selective coronary visualization. *AM HEART J* 83:360 1972

- 14 Nakhjavan F K Shedrovitzky H and Gold berg H Experimental myocardial infarction in dogs description of a closed chest technique *Circulation* 38:777 1968
- 15 Khomazyuk A I Nescheret A P and Kuz minsky N P Some new ways of experimental research of myocardial infarction *Kardiologiya* 5 (No 4) 19 1965
- 16 Ribellima J Selective embolization of the coronary arteries A hemodynamic metabolic and radiologic study *Proc Soc Exp Biol Med* 117:1367 1964
- 17 Johnsrude I S, and Goodrich J K An experimental partial occlusive device for vessels delivered by arterial catheter *Am HEART J* 77:805 1969
- 18 Gensini G G Palacio A Buonanno C Kelly A E and Muller W F Superselective coronary occlusion under cinefluorographic control in experimental animals technique and results (Abstr) *Circulation* 34:111 108 1966
- 19 Gensini G G Buonanno C Palacio A Kelly A E and Muller W F Cinefluorographic control of super selective coronary occlusion in experimental animals *J Soc Motion Picture and TV Engineers* 75 649 1966
- 20 Jacobey J A Taylor W J Smith G T Gorlin R and Harken D III A new therapeutic approach to acute coronary occlusion I Production of standardized coronary occlusion with microspheres *Am J Cardiol* 9:60 1962
- 21 Aggess C M Rosenberg M J Jacobs H I Binder M J Schneiderman A and Clark W G Prolonged shock in the closed chest dog following coronary embolization with graded microspheres *Am J Physiol* 170:536 1952
- 22 Palmer J D O'Rourke R A Olson M S and Pinchard R N Experimental myocardial infarction *Am HEART J* 81:729 1971
- 23 Guzman S V Swenson E and Mitchell R Mechanism of cardiogenic shock *Circ Res* 10:746 1962
- 24 Hammer J and Píša Z A method of isolated gradual occlusion of a main branch of a coronary artery in closed chest dogs *Am HEART J* 64:67 1962
- 25 Szamosi A Experimental occlusion of the coronary arteries in the closed chest dog: a selective method *Acta Radiol Diagn* 12 545 1972
- 26 Píša Z and Hammer J Experimental myocardial infarction in closed chest dogs with selective embolization of the coronary vascular bed, *Exp Med Surg* 19 1 1961
- 27 Herrmann G, and Decherd G Creatine mobilization in myocardial damage *Proc Soc Exp Biol Med* 32:177 1934
- 28 Lluch S Mogulevsky H C Pietra G Shaffer A B Hirsch, L J and Fishman A P A reproducible model of cardiogenic shock in the dog *Circulation* 39:205 1969
- 29 Lumicao II G Russell R O Jr and Rack ley C E Left ventricular performance in dogs with selective coronary embolization with mercury *Am J Med Sci* 261:27, 1971
- 30 Haft J I and Damato A N Measurement of collateral blood flow after myocardial infarction in the closed chest dog *Am HEART J* 77 641 1969
- 31 Rees J R and Redding V J Experimental myocardial infarction by a wedge method early changes in collateral flow *Cardiovasc Res* 2:43 1968
- 32 Weiss A II Senft A Khan M I and Regan T J Effect of nitrate infusions on the systemic and coronary circulations following acute experimental myocardial infarction in the intact dog *Am J Cardiol* 30 362 1972
- 33 Salazar A E Experimental myocardial infarction induction of coronary thrombosis in the intact closed chest dog *Circ Res* 9:1351 1961
- 34 David M S Charrette F J P and Lynn R B Experimental coronary artery thrombosis for production of cardiogenic shock *Can J Surg* 13:189 1970
- 35 Weiss A II Moschos C B Passanante A J Khan M I and Regan T J Relative effectiveness of three antiarrhythmic agents in the treatment of ventricular arrhythmias in experimental acute myocardial ischemia *Am HEART J* 81:503 1971
- 36 Cohen M V Downey J M Eldh P Urschel C W Sonnenblick E H and Kirk E S Enhancement of myocardial contractility by dilatation of coronary collaterals (Abstr) *Am J Cardiol* 31:126 1973
- 37 Schaper W The collateral circulation of the heart Amsterdam 1971 North Holland Publishing Co
- 38 Pasyk S Bloor C M Khouri E M and Gregg D E Systemic and coronary effects of coronary artery occlusion in the unanesthetized dog *Am J Physiol* 220 646 1971
- 39 Elliot E C Bloor C M Jones E L Mitchell W J and Gregg D E Effect of controlled coronary occlusion on collateral circulation in conscious dogs *Am J Physiol* 220:857 1971
- 40 Blum R L Alpern H Jaffe H Lang T W and Corday E Determination of interarterial coronary anastomosis by radioactive spheres Effect of coronary occlusion and hypoxemia *Am HEART J* 79:244 1970
- 41 Becker L C Fortuin N J and Pitt II Effect of ischemia and antianginal drugs on the distribution of radioactive microspheres in the canine left ventricle *Circ Res* 28:263 1971

Protection against epinephrine induced myocardial necrosis with clofibrate

Jacob I Haft MD

Paul D Kranz MD

Frank Albert MD

Rolf Oestreicher MD

Bronx NY

Recent clinical trials have suggested that clofibrate exerts a protective effect against myocardial infarction and death among middle aged men an effect that may be independent of the lowering of serum lipid levels by the drug.¹⁻³ Clofibrate has been demonstrated to decrease platelet adhesiveness among men with nonacute ischemic heart disease.⁴⁻⁶ It is possible that the precipitating event in acute myocardial infarction may be the sudden occlusion of a narrowed coronary artery by an intravascular platelet aggregate⁷ and it is possible that clofibrate may exert its protective effect by inhibiting such intravascular platelet aggregation.

Sympathetic catecholamines will cause platelet aggregation *in vitro*⁸⁻¹⁰ and infusion of norepinephrine will cause intravascular aggregation of platelets in the small vessels of the heart.¹¹ Rats subjected to stress will develop intravascular platelet aggregates in myocardial vessels.¹² Pretreatment of dogs with the antiplatelet aggregating drugs aspirin and dipyridamole will prevent the occurrence of the widespread myocardial necrosis usually seen after epinephrine

infusion.¹³ To determine if clofibrate will exert a similar protective effect against epinephrine induced cardiac necrosis suggesting that clofibrate might interfere with epinephrine induced platelet aggregation the following studies were performed.

Material and methods

Twenty two mongrel dogs weighing 15 to 20 kilograms were studied. Twelve dogs were pretreated for 5 days with clofibrate (Atromid S) 500 mg orally twice daily. Ten control dogs were not pretreated. Aortic pressure and the peripheral electrocardiogram were monitored as previously described.¹⁴

Epinephrine was infused intravenously at a rate of 4 µg per kilogram of body weight per minute for 4 hours. The dogs were killed 1 week after infusion the hearts were removed and fixed in formalin and histologic sections of the left ventricle were stained with hematoxylin and eosin and examined for presence or absence of necrosis. The degree of myocardial damage was graded on a scale of 0 to 3+ as previously described¹⁵ 3+ if extensive areas of con-

From the Cardiac Section of the Bronx Veterans Administration Hospital, and the Mount Sinai School of Medicine, New York, NY.

Received for publication March 19, 1973.

Reprint requests: Dr J I Haft, Chief Cardiac Section, Veterans Administration Hospital, 130 West Kingsbridge Rd, Bronx, NY 10468.

- 14 Nakhjavan F K, Shedrovitsky H and Goldberg H Experimental myocardial infarction in dogs: description of a closed chest technique *Circulation* 38:777 1968
- 15 Khomazuk A I, Nescheret, A P and Kuzminsky N P Some new ways of experimental research of myocardial infarction *Kardiologiya* 5 (No 4) 19 1965
- 16 Ribellima J Selective embolization of the coronary arteries: A hemodynamic, metabolic and radiologic study *Proc Soc Exp Biol Med* 117:367 1964
- 17 Johnsrude I S and Goodrich J K An experimental partial occlusive device for vessels delivered by arterial catheter *AM HEART J* 77:805 1969
- 18 Gensini G G, Palazzo A, Buonanno C, Kelly A E and Muller W F Superselective coronary occlusion under cinefluorographic control in experimental animals: technique and results (Abstr.) *Circulation* 34:III 108 1966
- 19 Gensini G G, Buonanno C, Palazzo A, Kelly A E and Muller W F Cinefluorographic control of super selective coronary occlusion in experimental animals *J Soc Motion Picture and TV Engineers* 75:649 1966
- 20 Jacobey J A, Taylor W J, Smith G T, Gorlin R and Harken D E A new therapeutic approach to acute coronary occlusion I: Production of standardized coronary occlusion with microspheres *Am J Cardiol* 9:60 1962
- 21 Agrest C M, Rosenberg M J, Jacobs H I, Binder M J, Schneiderman A and Clark W G Prolonged shock in the closed chest dog following coronary embolization with graded microspheres *Am J Physiol* 170:536 1952
- 22 Palmer J D, O'Rourke R A, Olson M S and Pinckard R N Experimental myocardial infarction *AM HEART J* 81:729 1971
- 23 Guzman S V, Swenson E and Mitchell R Mechanism of cardiogenic shock *Circ Res* 10:746 1962
- 24 Hammer J and Pisa Z A method of isolated gradual occlusion of a main branch of a coronary artery in closed chest dogs *AM HEART J* 64:67 1962
- 25 Szamosi A Experimental occlusion of the coronary arteries in the closed chest dog: a selective method *Acta Radiol Diagn* 12:545 1972
- 26 Pisa Z and Hammer J Experimental myocardial infarction in closed chest dogs with selective embolization of the coronary vascular bed, *Exp Med Surg* 19:1 1961
- 27 Herrmann G and Decherd G Creatine mobilization in myocardial damage *Proc Soc Exp Biol Med* 32:377 1934
- 28 Lluich S, Mogulovsky, H C, Pietra G, Shaffer A H, Hirsch, L J and Fishman A P A reproducible model of cardiogenic shock in the dog *Circulation* 39:205 1969
- 29 Lumicao B G, Russell R O Jr and Rackley C E Left ventricular performance in dogs with selective coronary embolization with mercury *Am J Med Sci* 261:27 1971
- 30 Haft J I and Damato A N Measurement of collateral blood flow after myocardial infarction in the closed-chest dog *AM HEART J* 77:641 1969
- 31 Rees J R and Redding V J Experimental myocardial infarction by a wedge method: early changes in collateral flow *Cardiovasc. Res.* 2:43 1968
- 32 Weiss A B, Senft A, Khan M I and Regan T J Effect of nitrate infusions on the systemic and coronary circulations following acute experimental myocardial infarction in the intact dog *Am J Cardiol* 30:362 1972
- 33 Salazar A E Experimental myocardial infarction: induction of coronary thrombosis in the intact closed chest dog *Circ. Res.* 9:1351 1961
- 34 David M S, Charrette E J P and Lynn R B Experimental coronary artery thrombosis for production of cardiogenic shock *Can J Surg* 13:189 1970
- 35 Weiss A B, Moschos C B, Passanante A J, Khan M I and Regan T J Relative effectiveness of three antiarrhythmic agents in the treatment of ventricular arrhythmias in experimental acute myocardial ischemia *AM HEART J* 81:503 1971
- 36 Cohen M V, Downey J M, Eldh P, Urschel C W, Sonnenblick E H and Kirk E S Enhancement of myocardial contractility by dilatation of coronary collaterals (Abstr.) *Am J Cardiol* 31:126 1973
- 37 Schaper W The collateral circulation of the heart *Amsterdam* 1971 North Holland Publishing Co
- 38 Paszyk S, Bloor C M, Khouri E M and Gregg D E Systemic and coronary effects of coronary artery occlusion in the unanesthetized dog *Am J Physiol* 220:646 1971
- 39 Elliot E C, Bloor C M, Jones E L, Mitchell W J and Gregg D E Effect of controlled coronary occlusion on collateral circulation in conscious dogs *Am J Physiol* 220:857 1971
- 40 Blum R L, Alpern H, Jaffe H, Lang T W and Corday E Determination of interarterial coronary anastomosis by radioactive spheres: Effect of coronary occlusion and hypoxemia *AM HEART J* 79:244 1970
- 41 Becker L C, Fortuin N J and Pitt B Effect of ischemia and antianginal drugs on the distribution of radioactive microspheres in the canine left ventricle *Circ Res* 28:763 1971

In another study from this laboratory it was shown that epinephrine induced myocardial necrosis can be prevented by pretreatment with drugs known to inhibit platelet aggregation.¹¹ In the present study we have demonstrated that epinephrine induced myocardial necrosis can be prevented similarly by pretreating with oral clofibrate a drug which has been described as possibly being effective in the long term treatment of ischemic heart disease in man. It is possible that clofibrate exerts its protective effects by an alteration of platelet function or of the response of platelets to exposure to sympathetic catecholamines.

Over the past decade a number of investigators have demonstrated effects of oral clofibrate on platelet function in patients with ischemic vascular disease. Platelet adhesiveness and platelet factor-3 activity are abnormally increased in patients with nonacute ischemic heart disease.^{12,17} The cause of these thrombogenic changes in platelet function is unknown. A number of investigators have indicated that these platelet function abnormalities are at least partially corrected by administration of clofibrate. Oral clofibrate administration has been found to significantly reduce both platelet adhesiveness and platelet aggregation in patients with nonacute atherosclerotic disease both when administered alone and when administered as Atromid, a combination of clofibrate with androstenedione.^{18,20,21} The androstenedione may or may not enhance this effect.^{18,21} Some investigators have observed that this effect of clofibrate on platelet adhesiveness and aggregation appears to be independent of changes in serum lipid levels.^{18,20} Human platelet factor-3 activity and availability may also be reduced by treatment with oral clofibrate.²⁰

Although most reports indicate that oral clofibrate interferes with platelet aggregation in man, this observation is not unanimous. One study of platelet adhesiveness was unable to confirm the reported effects of oral clofibrate,²² and another study found that the reduction of platelet adhesiveness was gradually lost after two months treatment.²³ Nevertheless the antiplatelet aggregating action of clofibrate indicated by most human studies and the similarity of the

results of the present dog study with those of the previous study which used known antiplatelet aggregators suggest that the present results can be explained by clofibrate's interference with platelet aggregation.

It is also possible that the protective action of clofibrate observed in dogs is due to interference with vascular effects of epinephrine (other than vasoconstriction) that may result secondarily in the formation of platelet aggregates. Shimamoto and colleagues^{24,25} have demonstrated that epinephrine will cause endothelial edema and release of a thromboplastic and platelet attracting substance in rabbit aorta. Thus it is possible that catecholamine induced platelet aggregation is mediated by endothelial damage and/or vascular release of a humoral platelet aggregator. Numerous studies have demonstrated the platelet aggregating tendency induced by other types of endothelial damage.^{26,27} Rowsell and associates²⁸ however have shown that the presence of a subjacent endothelium or vessel wall is not essential for the thrombogenic effect of epinephrine *in vivo*. Shimamoto's group^{24,25} has demonstrated that the platelet consuming effect of intravenous epinephrine can be blocked in man and in animals by pretreatment with a variety of unrelated compounds. It is possible that the protective effect of clofibrate observed in the present study might be due to a similar block of the vascular effects of epinephrine.

In addition to its effects on serum triglycerides and cholesterol, oral clofibrate also tends to reduce serum phospholipid levels.²⁰ Mustard and associates²⁹ have reported that alterations in the concentrations of various specific serum phospholipids can have varying effects on the tendency of platelets to aggregate. Thus it is conceivable that the protective effect of clofibrate observed in this study was mediated by alterations in the plasma lipid constituents.

The protection against epinephrine induced necrosis by pretreatment with clofibrate demonstrated in this study implies that the drug exerts a direct or indirect inhibiting effect on epinephrine induced intravascular platelet aggregation in dog hearts. Such an effect may have applica-

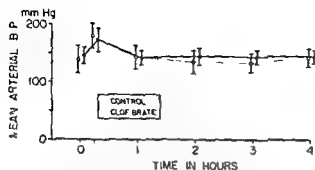


Fig. 1 The effect of epinephrine infusion on the mean arterial blood pressure of control dogs and those pretreated with clofibrate

fluent necrosis (destruction of myocardial cells presence of round cell infiltration and early fibrosis) were present, 2+ if extensive areas of necrosis without confluence were found, 1+ if small areas of necrosis more than 3 per slide were present on 6 slides and 0 if no areas of necrosis were found on any of the slides. Pretreated and control dogs were studied in random order, the grading pathologist (R. O.) did not know which or how many animals had been pretreated.

Results

The blood pressure responses to the epinephrine infusions were similar to those previously described by Moss and colleagues.¹⁶ After an early rise the pressure usually fell to near the control level at 1 hour and remained there for the duration of the infusion (Fig. 1). Systolic, diastolic, and mean pressures of the clofibrate group were not significantly different from those of the control group at comparable times during the infusions (Student *t* test).

On histologic study 1 week after infusion (Table I), all dogs in the control group had evidence of cardiac necrosis: 6 dogs with 3+, 2 dogs with 2+, and 2 dogs with 1+ necrosis. Among the 12 dogs pretreated with clofibrate 3 dogs had 2+ necrosis and 2 dogs had 1+ necrosis. Seven dogs had no evidence of necrosis. Comparison between the pretreated group and the control group with respect to the presence or absence of histologic evidence of necrosis showed that the findings were statistically significant ($p < 0.01$ by chi square analysis).

By the end of the infusion, few dogs had significant electrocardiographic changes.

Table I Histologic cardiac findings 1 week after infusion

| | Extent of necrosis* | | | |
|----------------------------|---------------------|----|----|----|
| | 3+ | 2+ | 1+ | 0+ |
| Control group (10 dogs) | 6 | 2 | 2 | — |
| Clofibrate group (12 dogs) | — | 3 | 2 | 7 |

*Necrosis is graded 1+ to 3+. Grade 0 indicates no evidence of necrosis. The difference between the 2 groups is statistically significant (see text).

One dog in the control group had an acute inferior wall myocardial infarction and 2 others had left axis shifts. These 3 dogs all had 3+ necrosis on histologic examination. One pretreated dog with 1+ necrosis had a right axis shift. All other dogs had either no electrocardiographic changes or only non-specific T wave changes.

Discussion

Production of diffuse patchy myocardial necrosis by catecholamine administration has been known since 1907.¹⁷ Since then this effect has been described following infusions of epinephrine, norepinephrine, or isoproterenol in a variety of animals and in man.^{18,19} Similar myocardial changes have been demonstrated in the majority of patients who die with pheochromocytoma.²⁰ In a study previously reported from this laboratory it was demonstrated that nor epinephrine infusions in doses known to cause such myocardial necrosis will result in intravascular platelet aggregation in the dog heart. Intravascular platelet aggregates were associated with changes of early myocardial necrosis before there was electron microscopic evidence of fibrinous thrombus formation.²¹ In 1967 Jorgensen and colleagues² demonstrated that intravascular platelet aggregates in the pig heart induced by intracoronary infusion of adenosine diphosphate will cause transient obstruction of the microcirculation leading to diffuse myocardial necrosis.²² Thus, it appears that myocardial necrosis can be induced directly by intravascular aggregation of platelets.

clofibrate and 10 dogs not pretreated were infused with epinephrine 4 µg per kilogram of body weight per minute. After killing the dogs 1 week later the degree of myocardial damage was assessed histologically. All control dogs had myocardial necrosis 6 with 3+ 2 with 2+ and 2 with 1+ necrosis. Seven of the 12 pretreated dogs had no necrosis 2 had 1+ and 3 had 2+ necrosis. These differences are statistically significant ($p < 0.01$). Previous studies have shown that norepinephrine infusions will induce intravascular platelet aggregation in the dog heart and pretreatment with antiplatelet aggregating agents will prevent epinephrine induced myocardial necrosis. The demonstration of oral clofibrate's protection against epinephrine induced myocardial necrosis suggests that clofibrate may exert a similar antiplatelet aggregating effect. It is possible that oral clofibrate exerts a beneficial effect in human coronary artery disease via a similar mechanism.

REFERENCES

- Dewar H A. Trial of clofibrate in the treatment of ischaemic heart disease. Five year study by a group of physicians of the Newcastle upon Tyne region. *Br Med J* 4:767 1971
- Oliver M F. Ischaemic heart disease: a secondary prevention trial using clofibrate. Report by a research committee of the Scottish Society of Physicians. *Br Med J* 4:775 1971
- Krasno L R and Kidera G F. Clofibrate in coronary heart disease. *J A M A* 219:845 1972
- Carson P, McDonald L, Pickard S, Pilkington T, Davies B and Love F. Effect of Atromid on platelet stickiness. *J Atherosclerosis Res* 3:619 1963
- Symons C de Toeseglin A and Cook I J Y. Effect of ethyl chlorophenoxyisobutyrate with or without androsterone on platelet stickiness. *Lancet* 2:233 1964
- Carson P, McDonald L, Pickard S, Pilkington T, Davies B and Love F. Effects of clofibrate with androsterone (Atromid) and without androsterone (Atromid S) on blood platelets and lipids in ischaemic heart disease. *Br Heart J* 28:400 1966
- Glynn M F, Murphy E A and Mustard J F. Effect of clofibrate on platelet economy in man. *Lancet* 2:447 1967
- Robinson R W and LeBeau R J. Platelet adhesiveness and aggregation with chlorophenoxyisobutyric ester. *Am J Med Sci* 253:76 1967
- Mustard J F and Packham M A. Platelet function and myocardial infarction. *Circulation* 40 (Suppl 4):20 1969
- O'Brien J R. Some effects of adrenaline and anti-adrenaline compounds on platelets in vitro and in vivo. *Nature* 200:763 1963
- O'Brien J R. Variability in the aggregation of human platelets by adrenaline. *Nature* 202:1188 1964
- Mitchell J H and Sharp A A. Platelet clumping in vitro. *Br J Haematol* 10:78 1964
- Haft J I, Kranz P D, Albert F J and Fani K. Intravascular platelet aggregation in the heart induced by norepinephrine. Microscopic studies. *Circulation* 46:698 1972
- Haft J I and Fani K. Intravascular platelet aggregation in the heart induced by stress. *Circulation* 47:353 1973
- Haft J I, Gershengorn K, Kranz P D and Oestreicher H. Protection against epinephrine-induced myocardial necrosis by drugs that inhibit platelet aggregation. *Am J Cardiol* 30:838 1972
- Moss A J, Vitands I and Schenk E. Cardiovascular effects of sustained norepinephrine infusions. *Circ Res* 18:596 1966
- Josue O. Hypertrophie cardiaque causee par l'adrenaline et la toxine typhique. *Compt Rend Soc Biol (Paris)* 63:285 1907
- Szalacs J E and Mehlman B. Pathologic changes induced by l-norepinephrine. *Am J Cardiol* 5:619 1960
- Maling H M, Highman B and Thompson E C. Some similar effects after large doses of catecholamines and myocardial infarction in dogs. *Am J Cardiol* 5:628 1960
- Raab W, Stark E, MacMillan W H and Gigue W R. Sympathogenic origin and anti-adrenergic prevention of stress induced myocardial lesions. *Am J Cardiol* 8:203 1961
- Wexler B C, Judd J T and Kittinger G W. Myocardial necrosis induced by isoproterenol in rats. *Angiology* 19:665 1968
- Ferrans V J, Hibbs R G, Black W C and Weilbaecher W G. Isoproterenol induced myocardial necrosis. *Am Heart J* 68:71 1964
- Schenk E A and Moss A J. Cardiovascular effects of sustained norepinephrine infusions. 2. Morphology. *Circ Res* 18:605 1966
- Van Vleet P D, Burchell H H and Titus J L. Focal myocarditis associated with pheochromocytoma. *N Engl J Med* 274:1102 1966
- Jorgensen L, Rowsell H C, Hovig T, Glynn M F and Mustard J F. Adenosine diphosphate induced platelet aggregation and myocardial infarction in swine. *Lab Invest* 17:616 1967
- McDonald L and Edgill M. Coagulability of the blood in ischaemic heart disease. *Lancet* 2:457 1957
- Norday A and Rodset J M. Platelet phospholipids and their function in patients with ischaemic heart disease. *Acta Med Scand* 183:133 1970
- Clibert J B and Mustard J F. Some effects of Atromid on platelet economy and blood coagulation in man. *J Atherosclerosis Res* 3:623 1963
- Gibelli A, Frandoli G, Girola P and De Nicola P. Fibrinolytic activity and reduction

tions to human disease. Acute myocardial infarction has been described to occur commonly following prolonged emotional stress.^{11,12} During stress the endogenous secretion of sympathetic catecholamines is increased in man.¹³ This laboratory has recently reported the occurrence of extensive intravascular platelet aggregation in the hearts of rats sacrificed following a period of prolonged physical and emotional stress.¹⁴ It is possible that the higher levels of circulating catecholamines associated with human emotional stress in combination with pre-existing atheromatosis may lead to an enhanced tendency to form intravascular platelet aggregates and platelet thrombi. If one of these thrombi forms at or embolizes to an area of a coronary artery already narrowed by an atherosclerotic plaque, occlusion of the lumen and ischemia or infarction may ensue. Platelet thrombi are fragile and hence, typically, such an infarction would not necessarily be associated with identifiable thrombosis on postmortem examination¹⁵ and thus would be consistent with the low pathological incidence of coronary thrombosis reported in acute fatal myocardial infarction.¹⁶ Therefore one might anticipate that pharmacologic agents which inhibit catecholamine-induced platelet aggregation would be effective in reducing the incidence of ischemic episodes in patients with coronary artery disease subject to emotional stress.

In this regard it is noteworthy that three different long term studies recently have reported that oral clofibrate reduces the incidence of myocardial infarction in patients with and without coronary artery disease. It is claimed that clofibrate's protective action in these patients bears little relation to its lipid lowering effect. Krishna and Kalaria¹ reported 518 men treated with 500 mg of clofibrate 4 times daily for 39 months and compared them to 550 male controls pair matched for risk factors of coronary artery disease. Ninety eight per cent of their patients were free of signs or symptoms of coronary artery disease at the start of observation. The incidence of episodes of angina pectoris and myocardial infarction were monitored in both groups and were found to be significantly higher in the control group than in the treated

group. These differences were significant to the 0.1 per cent level ($p < .001$). Those hyperlipidemic men treated with clofibrate who failed to sustain a reduction of their serum lipid levels nevertheless demonstrated no clinical or electrocardiographic evidence of myocardial infarction in contrast to a 58 per cent incidence of infarction among the hyperlipidemic controls. In the United Kingdom, physicians of the Newcastle upon Tyne region¹ reported a 5 year double blind study of 244 patients with ischemic heart disease treated with clofibrate 1.5 to 2.0 Gm daily and compared them with 253 patients treated with placebo. The incidence of cardiac deaths was significantly lower in the treated group ($p = .002$) as was the incidence of first nonfatal myocardial infarction ($p < .005$). Those male patients who sustained cardiac death or infarction had had at least an inadequate reduction of serum lipid levels in response to treatment as those who had no such incidents. The Scottish Society of Physicians² also reported a 6 year study of patients with pre-existing coronary artery disease. Three hundred and fifty patients were treated with clofibrate 1.6 to 2.0 Gm daily and were compared with 367 similar patients treated with placebo. Clofibrate significantly reduced the incidence of cardiac death in patients entering the study with a history of angina pectoris ($p < .001$) and the incidence of myocardial infarction in patients entering with a history of both angina and previous infarction ($p < .005$). There was no significant relationship between the response of serum cholesterol to clofibrate and the incidence of events either in patients with angina or in those with infarction.

It is quite possible that the beneficial effects of oral clofibrate noted in all these three studies are related to an inhibitory effect upon intravascular platelet aggregation. It is hoped that further clinical trials using known inhibitors of platelet aggregation will be forthcoming.

Summary

To investigate the possibility that oral clofibrate interferes with catecholamine-induced platelet aggregation and myocardial necrosis 12 dogs pretreated with oral

Congenital absence of the circumflex coronary artery

Clinical and cinearteriographic observations

Vincent Barrest MD

Armando Susmano MD

Michael A Colandrea MD

Maurice L Bogdonoff MD

Joseph J Muenster MD

Chicago Ill

The current emphasis on the surgical management of occlusive coronary artery disease has necessitated a deeper understanding of the anatomy of the coronary vessels. Precise knowledge of this anatomy becomes critical in the preoperative interpretation of coronary arteriograms. Congenital coronary anomalies per se have received modest attention in the clinical literature with the exception of coronary vessels arising from the pulmonary artery.

The variations in normal coronary anatomy and the specific congenital anomalies have been catalogued by a number of authors.^{1,2,3,4} Blake and colleagues² have grouped all anomalies into three categories which are based on the degree of impaired function encountered and the extent of the technical impediment for the cardiovascular surgeon.

The most commonly described variation of the left circumflex artery occurs in 1 to 2 per cent of cases in which it originates from

the proximal portion of the right coronary artery or from the right coronary cusp.^{1,2,3,4} No reference is made to complete absence of the left circumflex artery.

This report presents two cases which we believe represent congenital absence of the left circumflex artery with all other vessels being normal.

Case reports

Case 1 Mr M M is a 33 year old Caucasian machanicist who was referred to Rush Presbyterian-St. Luke's Medical Center for evaluation of coronary artery disease. He was in apparent good health until March 1972 at which time he developed substernal non radiating dull chest pain while sitting at home. The pain lasted for approximately five minutes and was associated with no other symptoms. The following day the pain recurred and was associated with some shortness of breath. No specific measures were taken and he recovered. However in April 1972 the substernal discomfort recurred and was sufficiently severe that he was admitted to a local hospital where an electrocardiogram was read as showing an anteroseptal myocardial infarction. Pain recurred in May and September 1972 and the patient was hospitalized on each occasion. The elec-

From the Section of Cardio Respiratory Diseases, Department of Medicine and Department of Radiology Rush Presbyterian-St. Luke's Medical Center and Rush Medical College Chicago Ill.

This work was supported in part by United States Public Health Service Training Grant N 5 TO 1 HL-05714-07 from the National Heart and Lung Institute of the National Institute of Health.

Received for publication April 27 1973

Reprint requests to Armando Susmano MD Rush Presbyterian-St. Luke's Medical Center 1753 W. 4th St. Chicago 9 Parkway Chicago, Ill. 60612

- of platelet adhesiveness following prolonged administration of ethyl chlorophenoxy isobutyrate (Atromid S) *Hemostase* 6:285 1966
- 30 Norday A and Rodset J M Platelet function and platelet phospholipids in patients with hyperbetalipoproteinemia Effect of nicotinic acid and clofibrate, *Acta Med Scand* 189:385 1971
- 31 O'Brien J R and Heywood J H A comparison of platelet stickiness test during an Atromid S trial *Thromb Diath Haemorrh* 16:768 1966
- 32 Chakrabarti R Fearnley G H and Evans J F Effects of clofibrate on fibrinolysis platelet stickiness plasma fibrogen and serum cholesterol *Lancet* 2:1007 1968
- 33 Shimamoto T and Osjoeka T Release of a thromboplastic substance from arterial walls by epinephrine *Circ Res* 12:138 1963
- 34 Shimamoto T The relationship of edematous reaction in arteries to atherosclerosis and thrombosis *J Atherosclerosis Res* 3:187 1963
- 35 Honour A J and Ross Russell R W Experimental platelet embolism *Br J Exp Pathol* 43:350 1962
- 36 Honour A J Mitchell J R A Platelet clumping in vivo *Nature* 197:1019 1963
- 37 Born G V R Honour A J and Mitchell J R A Inhibition by adenosine and by 2-chloroadenosine of the formation and embolization of platelet thrombi *Nature* 202:761 1964
- 38 Rowse H C Hegardt B Downie H G Mustard J F and Murphy E A Adrenaline and experimental thrombosis *Br J Haematol* 12 66 1966
- 39 Yamazaki H Sano T Odakura T Takenchi K Matsumura T Hosaki S and Shimamoto T Appearance of thrombogenic tendency induced by adrenaline and its prevention by β adrenergic blocking agent nialamide and pyridinol carbamate *Thromb Diath Haemorrh* 26 251 1971
- 40 Hellman L Zumoff B Kessler G Kara E Rubin I L and Rosenfeld R S Reduction of serum cholesterol and lipids by ethyl chlorophenoxyisobutyrate *J Atherosclerosis Res* 3:154 1963
- 41 Mustard J F Packham M A Rowse H C and Jorgensen L The role of thrombogenic factors in atherosclerosis *Ann N Y Acad Sci* 119 848 1968
- 42 Dreyfuss F Role of emotional stress preceding coronary occlusion *Am J Cardiol* 3 590 1959
- 43 Wolf S Psychosocial forces in myocardial infarction and sudden death *Circulation* 40 (Suppl 4) 74 1969
- 44 Nestel P J Vergheze A and Lovell R R H Catecholamine secretion and sympathetic nervous responses to emotion in men with and without angina pectoris *Am Heart J* 73:227 1967
- 45 Ehrlich J C and Shinohara Y Low incidence of coronary thrombosis in myocardial infarction *Arch Pathol* 78:432 1964



Fig 2 Lateral view of the right coronary artery which is entirely normal and supplies long branches to the posterior wall



Fig 3 Lateral projection showing a double platinum pacing catheter in the right ventricular chamber. The main left coronary artery is visualized. The ostium of the vessel is posteriorly located. The vessel is continuous with the left anterior descending artery which appears to be entirely normal. A true main circumflex artery is not visualized.

35 mm film was exposed at 60 frames per second with kilovoltage and milliamperage automatically regulated. The film was processed in a Kodak Versamat film processor.

The right coronary artery was entirely free of occlusive mural disease and was shown to supply a large portion of the posterior surface of the heart (Fig 2). The main left coronary artery demonstrated a more posterior origin and continued as the anterior descending artery which was normal. No charac-

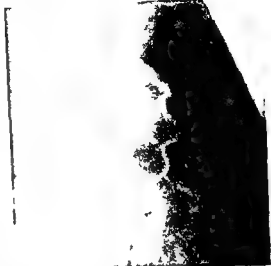


Fig 4A Anteroposterior projection of the main left coronary artery injection. The first branch of the left coronary artery is seen. This small vessel courses laterally in the usual area of distribution of the left anterior descending.

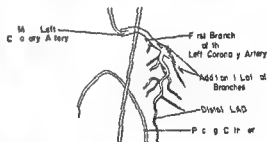


Fig 4B Detailed illustration outlining the various branches of the left coronary artery as shown in Fig 4A.

teristic circumflex branch could be identified (Fig 3). Multiple selective catheter injections were performed and no accessory ostia were visualized. The earliest branches of the left anterior descending artery did not follow the typical course of the main circumflex artery when seen in the posteroanterior projection (Fig 4A and B).

Case 2: Mr. D. M. is a 50-year-old Caucasian man, production engineer with a five-year history of crampy precordial pain radiating to the left arm. In 1968 he was hospitalized for 21 days for suspected myocardial infarction with no concrete evidence to support that diagnosis. Beginning in March 1972 he began to notice more frequent episodes of chest pain as often as two to three times daily. These episodes were occasionally associated with nausea, diaphoresis, and dizziness, however, there was equivocal response to nitroglycerine, isosorbide dinitrate. He had a history of mild hypertension for approximately five years and greater than 30-pack years of cigarette

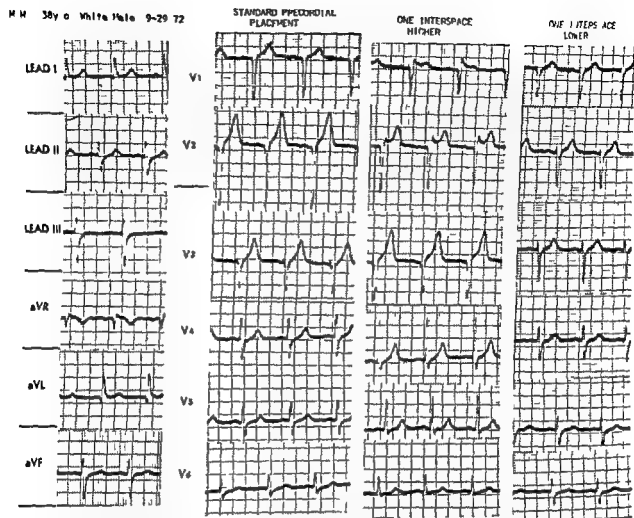


Fig 1 A 12 lead electrocardiogram and six precordial leads taken at three different intercostal levels (Case 1) A normal progression of the R wave can be seen when the electrodes are placed one interspace below a standard location

trocardiograms were again interpreted as showing an anteroseptal myocardial infarction. During these hospitalizations there were no enzyme changes and the patient had an inconsistent response to the administration of sublingual nitroglycerine. Complete upper gastrointestinal and gall bladder radiologic survey were normal. The physical examination was consistently within normal limits.

The patient's father had maturity onset diabetes adequately controlled with diet. There are no other significant features in the patient's family history. The only other pertinent risk factor for coronary disease in this patient was a history of 40 pack years of cigarette smoking.

On admission to the Rush Presbyterian-St. Luke's Medical Center the physical examination was entirely normal. Hemoglobin was 14.2 Gm per cent and complete blood count was normal. The two-hour postprandial blood sugar was 100 mg per cent. Uric acid, serum cholesterol and triglyceride levels were normal. The posteroanterior and left lateral chest x ray were also normal. A 12 lead electrocardiogram at rest was within normal limits. Precordial leads were taken at the standard level, at one interspace higher and at one interspace lower (Fig

1). The high precordial sequence revealed a QS configuration in V_1 through V_2 with upward scooping of the ST segment in V_2 , which had been interpreted earlier as consistent with anteroseptal myocardial infarction. However, the precordial sequence at the standard level showed an entirely normal progression of the R wave with concordant T waves. The QR-ST angle in the frontal plane was normal with a mean frontal QRS axis of -30 degrees. The patient underwent a 44 step double Master's exercise test which revealed no changes. A vectorcardiogram performed on a Hewlett Packard (Model 1507A) vector programmer using the Frank lead system was interpreted as within normal limits.

Selective coronary arteriography and left ventriculography were performed by the Seldinger method using Judkins Cordis catheters.* Simultaneous biplane cinearteriography was performed employing Siemens† cine pulsed equipment with high resolution cesium iodide phosphor Kodak‡

*Cordis Corporation Miami Fla.

†Siemens Corporation biplane X ray units III mm variable speed Model BS100 Erlangen Germany

‡Kodak Linagraft Shelburne Ester A11 Base Type 2476 Oakbrook, Ill

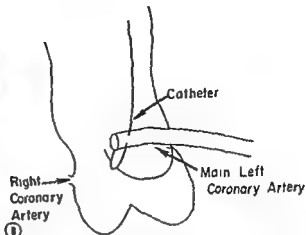


Fig 8A and B A Aortic root opacification showing a normal aortic valve and no accessory coronary ostia. Again the main left artery gives off no circumflex branch and is continuous with a normal left anterior descending artery. B Detailed illustration outlining the anatomy of the aortic cusp and its relationship to a single ostium of the left coronary artery.

cardiogram were normal. Treadmill exercise was attempted using a standard protocol. After 1 minute at 0 per cent incline and 3 m.p.h. the patient began to experience arm pain and the exercise was interrupted. The heart rate at rest was 50 beats per minute and was increased to 68 with no evidence of myocardial ischemia at this point. A vectorcardiogram was also performed which was normal (Fig 5).

Coronary arteriography was performed. The main left coronary artery originated more posteriorly than usual and continued as the left anterior descending which was entirely normal. No circumflex artery was identified (Fig 6). Selective left cusp injections (Fig 7) and an aortic root angiogram (Fig 8A and B) demonstrated the unusual origin of the proximal vessel and the lack of accessory ostia. The right coronary artery injection revealed a normal vessel with prominent posterior distribution (Fig 9).



Fig 9 Lateral view of the right coronary artery showing a markedly dominant distribution. Reflux of dye into the cusp suggests no accessory ostia. The vessel is entirely normal.

Discussion

The incidence of anomalous branches of the left coronary artery is relatively rare. In his compendium, James⁵ indicates that the only consistently occurring anomaly of the left circumflex is that in which the vessel originates from the main right coronary artery or from an accessory ostium in the right coronary cusp, the former being more frequent. In this setting, the vessel travels posteriorly around the aorta to enter the atrioventricular groove.^{2,3,7,8} The defect is particularly frequent in patients with transposition of the great vessels.⁷ Our angiograms clearly exclude this possibility.

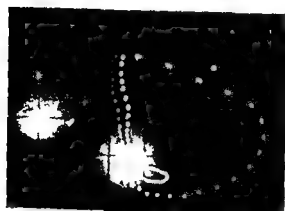
Wenger and Peace¹² and Grossman and Adams⁴ have reported cases of the diminutive left coronary syndrome. In these pa-

tients the circumflex artery is absent or is an insignificant vessel, but the left anterior descending artery is also abnormal. Again our angiograms exclude this possibility.

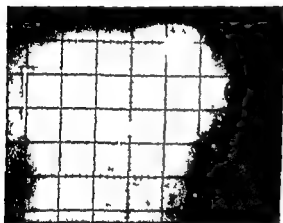
The selective cusp injections in Case 1 and the aortic root injection in Case 2 effectively exclude the possibility of accessory origin of the left circumflex from the left coronary cusp, which is seen in 1 per cent of cases.¹³

In Case 2, total occlusion of the circumflex artery at the ostium seems unlikely in

7' 1", 51 y o White male 8-37-72



HORIZONTAL PLANE



FRONTAL PLANE



RIGHT SAGITTAL

Fig 5 Display of a normal vectorcardiogram obtained in Case 2

smoking. There was no evidence of diabetes mellitus or gout but the family history was impressive. The patient's father suffered three cerebrovascular accidents and his mother died at the age of 42 from cerebrovascular disease. Two sisters and one brother have all had documented myocardial infarctions.

The patient was admitted to the Rush Presbyterian-St. Luke's Medical Center for further evaluation of his chest pain. On admission the physical examination was completely normal. Hemoglobin



Fig 6 Lateral projection showing a bipolar pacing wire in the right ventricular chamber. The main left coronary artery originates from an unusual posterior location in the left coronary cusp and is continuous with an entirely normal left anterior descending artery. There is no identifiable main circumflex artery or ramus intermedius. The posterior wall of the aorta and the coronary cusp is opacified by refluxed dye and no accessory ostia are visualized.

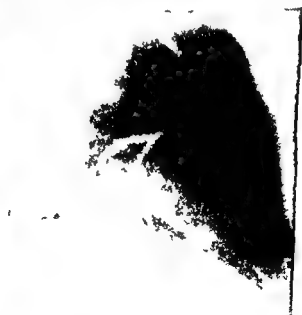


Fig 7 Lateral projection of a selective left coronary cusp injection showing no circumflex branch of the left coronary artery. The main left coronary has an unusual posterior origin in the left coronary cusp.

complete blood count, serum electrolytes, and enzyme levels were all normal. Liver function studies, serum cholesterol, and serum triglycerides were also normal. The upper gastrointestinal evaluation was normal. Routine chest x-ray and 12 lead electro-

Traumatic coronary arterial fistula

A case report and review of the literature

Richard R. Liberthson, M.D.

Kenneth Barron, M.D.

J. Warren Harthorne, M.D.

Robert E. Dinsmore, M.D.

Willard M. Daggett, M.D.*

Boston, Mass.

Traumatic coronary artery fistulas are rare. Ten cases have been reported previously¹⁻¹¹ (See Table 1). These cases elucidate the spectrum of problems involved in managing this entity. The present report discusses an additional case, summarizes the accumulated experience from the literature, and emphasizes some of the more difficult management problems associated with traumatic coronary arterial fistula.

Case report

A 24-year-old male heroin addict was shot at close range with a .22-caliber bullet on July 18, 1972. He was taken to a suburban hospital where his vital signs were stable and chest x-ray essentially normal except for the bullet, which was shown to be outside the chest wall beneath the right pectoralis muscles. No heart murmur was initially heard. An entrance and exit wound of the left biceps and an entrance wound in the left chest were observed. On the third day he became febrile, accumulated a marked right and left hemothorax, and was noted for the first time to have a loud precordial murmur. The patient was transferred to the Massachusetts

General Hospital on July 21, 1972, where he presented with a blood pressure of 120/70 mm Hg, temperature of 101.1°, heart rate of 150 per minute, and respiration of 40 per minute. He was pale and dyspneic. The heart was mildly enlarged, a summation gallop was present, and a loud high frequency continuous murmur was heard over the entire precordium.

The ECG showed borderline LVH and diffuse changes consistent with acute pericarditis. An aortic root and left ventricular angiogram on July 21, 1972, revealed a shunt from the left anterior descending (LAD) coronary artery to the right ventricle (RV), an intact ventricular septum, and normal great vessels. The hemothorax was treated by closed thoracostomy drainage, and heroin withdrawal was begun. No symptoms ensued from the fistula. The murmur persisted, and the ECG evolved anterolateral ischemic changes. Cardiac catheterization on August 1, 1972, showed a negligible RV O₂ stepup and a mildly elevated LVEDP. Selective coronary angiography demonstrated a large LAD to RV shunt with diminished LAD flow distal to the fistula (Fig. 1). Because of persisting ischemia and suspected impending infarction, surgical correction of the coronary artery steal was performed on August 6, 1972. An entrance wound was observed just beneath the LAD about 4 cm from its origin; the myocardial wound of exit was on the diaphragmatic

From the Cardiac and Cardiovascular Surgical Unit and the Department of Medicine, Radiology, and Surgery, Massachusetts General Hospital and Harvard Medical School, Boston, Mass.

Supported in part by the following United States Public Health Service Grants: HL-06664 and HL-5196.

Received for publication May 3, 1973.

Reprint requests: Willard M. Daggett, M.D., Department of Surgery, Massachusetts General Hospital, Boston, Mass. 02114.

Dr. Daggett is an Established Investigator of the American Heart Association.

81 December 1973

American Heart Association

view of the absence of any demonstrable mural or occlusive disease of other vessels. In Case 1, one could argue that the earliest branches of the left coronary, in fact, represent a diminutive circumflex system in a patient with a massively right dominant system. Ninety per cent of patients will have this distribution of vessels,^{1,2} but most often a well defined circumflex vessel enters the atrioventricular groove. In the cases reviewed by James⁴ only 1 per cent of the vessels terminated before the crux. In our patient the posteroanterior projection of this vessel is atypical for a circumflex artery (Fig 4A and B). Further, the initial branch in question when viewed in motion, moves synchronously with all other branches of the left anterior descending, confirming its lateral course. Again, none of the vessels visualized shows any evidence of disease so that the likelihood of a totally occluded vessel is slight, at best.

Selective metabolic studies sampling the coronary sinus were not performed; therefore, we are unable strictly to exclude the possibility of myocardial ischemia.

Patient M. M. (Case 1) had a history of hospitalization for psychoneurosis 10 years prior to his cardiac evaluation. Since coronary arteriography, his pain has markedly diminished without the use of cardiac medications. His physical activities are unrestricted.

Patient D. M. (Case 2) has continued to have precordial pain, but there has never been electrocardiographic confirmation of ischemia. His daily activities have been voluntarily curtailed to exclude excessive exertion.

We believe, therefore, that the angiograms in these two cases represent congenital absence of the circumflex artery associated with no specifically abnormal electrocardiographic pattern.

Summary

Two patients found to have congenital absence of the circumflex coronary artery are reported. The clinical, electrocardiographic, vectorcardiographic, and cine angiographic findings are presented. No distinctive clinical or laboratory abnormalities were found except for the unusual anatomical posterior origin of the ostium of the main left coronary artery.

REFERENCES

- 1 Baroldi G and Scmazzone G. Coronary circulation in the normal and the pathologic heart. AFIP Monograph U. S. Govt. Printing Office 1967.
- 2 Blake H A et al. Coronary artery anomalies. *Circulation* 30:927 1964.
- 3 Dadds J V. Congenital anomaly of the left circumflex coronary artery. *J Pathol Bacteriol* 79:204 1960.
- 4 Grossman L A and Adams C W. Diminutive coronary artery syndrome. *JAMA* 188:1111 1964.
- 5 James T N. Anatomy of the coronary arteries. New York 1961. Paul B Hoeber Inc.
- 6 Judkins M P. Percutaneous transfemoral selective coronary arteriography. *Radiol Clin North Am* 6:467 1968.
- 7 Nora J J and McNamara D G. Anomalies of the coronary arteries and coronary artery fistula. In Watson H ed. *Pediatric cardiology*. St. Louis 1968. The C V Mosby Company.
- 8 Ogden J A. Congenital variations of the coronary arteries. Thesis 1968. School of Medicine Yale University.
- 9 Ogden J A. Congenital anomalies of the coronary artery. *Am J Cardiol* 25:474 1970.
- 10 Smith G T. The anatomy of the coronary circulation. *Am J Cardiol* 9:327 1962.
- 11 Smith J C. Review of single coronary artery with report of two cases. *Circulation* 11:68 1950.
- 12 Wenger N K and Peace R J. Rudimentary left coronary artery. *Am J Cardiol* 8:519 1961.
- 13 Zumbo O et al. Coronary atherosclerosis and myocardial infarction in hearts with anomalous coronary arteries (Abstr. from International Academy of Pathology). *Lab Invest* 14:571 1965.

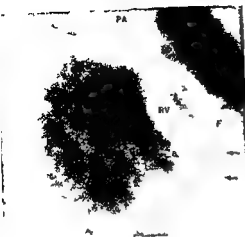


Fig 1 Selective injection of the left coronary artery shows the dilated LAD coronary artery (arrows) the site of the fistula (F) and flow of dye through the fistula into the RV. The angiogram also documents a coronary steal with markedly diminished flow into the distal LAD.



Fig 2 Operative photograph showing the site of the fistula in the back wall of the LAD which has been opened on its anterior surface. Proximal and distal control of the LAD and a large diagonal branch has been obtained by elastomer tapes. The entire heart is edematous.

present or detected at the time of initial injury and in several were not manifest until years later. Again as illustrated by our case symptoms were frequently not related to the fistula. In most patients a hemothorax was present and several were admitted in overt shock. In all instances initial wound recovery was uneventful.

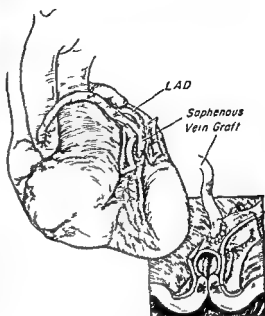


Fig 3 Diagrammatic representation of the surgical repair of the fistula between the LAD coronary artery and the RV. The fistula in the back wall of the LAD was closed by using mattress sutures placed behind the LAD. These sutures were buttressed by strips of Teflon felt. Flow to the distal LAD was assured by a complementary saphenous vein bypass graft inserted into the LAD just beyond the fistula.

Acute myocardial infarction was present in five cases and ours and one other had changes of acute myocardial ischemia. These changes were in the distribution of the involved coronary artery. Patients may be asymptomatic or have minimal symptoms related to these fistulas as in two patients followed for 9 and 20 years. In one case however progressive congestive heart failure necessitated surgery after eight months in two others angina pectoris developed after 4 and 12 years necessitating surgery and in a third ischemia with exercise led to correction after three years. Eight cases had operative correction ranging from three weeks to 12 years after injury. The indication cited most frequently was a dilated or enlarging coronary artery proximal to the fistula with concern for aneurysm progression or rupture. Since pericarditis is a frequent accompaniment of these lesions along with actual myocardial trauma ECC and enzyme measurements may be unreliable in assessing the

Table 1 Traumatic coronary artery fistulas reported in the literature

| Case | Source | Age (yr) | Type of wound | Site of fistula | Management | Results and follow-up |
|------|---|----------|------------------------------|---------------------------------|------------------------|--|
| 1 | Parmley 1958 Bravo 1971 (20-year follow up) Jaffee 1973 | 20 | Shrapnel | RC to RA | Observation | Mild dyspnea with exertion after 20 years |
| 2 | Jones 1966 | 18 | Gunshot | RC to RA and RV | Surgery after 4 months | Murmur recurred 3 weeks after surgery patient asymptomatic after 3 months |
| 3 | Tsagaris 1966 | 33 | Shrapnel | RC to RA | Surgery after 12 years | Murmur recurred one day after surgery and re-operated |
| 4 | Storey 1967 | 17 | Gunshot | RC to RA | Surgery after 9 years | Asymptomatic after 7 years |
| 5 | Serot 1968 | 30 | Stab wound | RC to right coronary vein to RA | Surgery after 4 months | Murmur recurred within hours of surgery and re-operated asymptomatic after 2 years |
| 6 | North 1971 | 19 | Gunshot | RC to RA | Surgery after 6 months | Asymptomatic |
| 7 | Lynn 1971 | 21 | Contusion from auto accident | RC to RV | Surgery after 8 months | Murmur recurred after 6 weeks and re-operated normal examination after 5 years |
| 8 | Morgan 1972 Kahn 1967 (2 year follow up) Forker 1971 (6-year follow up) | 17 | Contusion from auto accident | LAD to RV | Surgery after 3 years | Murmur recurred after 3 months |
| 9 | Constantino 1972 | 37 | Gunshot | LAD to RV | Observation | Asymptomatic |
| 10 | Tseng 1973 | 38 | Stab wound | LAD to LV | Surgery after 4 years | Apical pancytotoxic murmur of undetermined etiology after surgery asymptomatic |

surface of the RV. The region of the LAD was markedly edematous and the myocardium of the anterolateral aspect of the left ventricle was soft and boggy. A continuous thrill was palpable over the dilated proximal LAD. With the aid of cardiopulmonary bypass a 4 mm tear in the posterior surface of the LAD was exposed by a longitudinal arteriotomy in the LAD. This opening communicated via a fistulous channel to the RV near the origin of its outflow (Fig 2). The LAD and adjacent fistula were under tension with silk mattress sutures buttressed by strips of Teflon felt. A saphenous vein bypass graft from the aorta to the distal LAD was placed to insure adequate circulation to the anterolateral left ventricular wall (Fig 3).

The patient recovered uneventfully but on the fifth postoperative day a similar but softer continuous precordial murmur was again heard. A repeat coronary angiogram on August 16, 1972 revealed a patent bypass graft and LAD and a diminutive shunt from the LAD to the RV at the site of the initial fistula. Postoperative ECGs showed only nonspecific ST-T wave changes. It was decided not to attempt further correction of the persisting small shunt. When seen in follow-up after six months the patient was entirely asymptomatic. Physical examination revealed only a Grade I/VI ejection systolic murmur and the ECG was normal.

Discussion

Although congenital coronary artery fistulas are not uncommon, fistulas secondary to trauma are rare. The present case brings the total number reported to eleven. It also illustrates the remarkable homogeneity of this entity. The mean age of the reported cases is 25 years. Four were caused by bullet wounds, two by contusion incurred in automobile accidents, two by shrapnel and two by knife wounds. Right coronary (RC) to right atrial (RA) fistulas were most frequent. Our case and two others had LAD to RV shunts and one had an LAD to left ventricular (LV) communication. Aside from the fistulas, two cases had RA lacerations which required immediate repair, one had an LV laceration, one had a VSD and one a torn tricuspid leaflet. Otherwise associated cardiac damage was minimal.

Loud continuous precordial murmurs were the hallmark of these lesions. As in the present case, murmurs were not always

seven cases with unusual features. *Am J Cardiol* 30:432, 1972.

- 12 Jaffee, R. B., Glancy, L., Epstein, S. E., Brown, B. G. and Morrow, A. G. Coronary arterial right heart fistulae. Long term observations in seven patients. *Circulation* 47:133, 1973.
- 13 Constantino, T., Falcone, M. W., Perloff, J. K., DeLeon, A. C. Jr. and Jamison, W. L. Trau-

matic coronary arterial right ventricular fistula.

Report of a case. *Med Ann D C* 41:70, 1972.

- 14 Cheng, T. O. and Adkins, P. C. Traumatic aneurysm of left anterior descending coronary artery with fistulous opening into left ventricle and left ventricular aneurysm after stab wound of chest. Report of case with successful surgical repair. *Am J Cardiol* 31:384, 1973.

presence of myocardial ischemia. In our case, the primary indication for surgery was correction of the angiographically demonstrated coronary steal phenomenon, relief of distal ischemia evidenced by diminished LAD blood flow, and prevention of actual infarction. The present case was the only one in which a saphenous vein bypass was used. Follow up is too short to evaluate prevention of infarction but the frankly ischemic preoperative electrocardiogram reverted to only nonspecific changes at the time of hospital discharge and was normal at six months. Endocarditis was an additional concern, particularly with the associated heroin addiction. In none of the reported cases, however, even those with very long intervals prior to correction, was endocarditis reported although this may be due to prophylaxis in some. In five corrected patients, the continuous murmur recurred after surgery and required a second operation and in one other a new presystolic murmur of undetermined etiology was reported. In the present case the recurrent shunt was so small that repeat closure was not recommended. The frequency of recurrence highlights some of the surgical difficulties. Early correction unless necessitated by acute circumstances or, as in our case to prevent potential infarction, should best be delayed to allow subsidence of edema and swelling, thereby permitting better surgical delineation, correction, and healing. It is likely in our case that recurrence of the fistula was due either to opening of a covert channel as myocardial edema resolved or development of a new fistula secondary to postoperative tissue necrosis. The reported patients show recurrence developing up to three months after surgery. Most reported corrections were remote in time from the initial injury so inadequate healing is not the only factor. Reconnection and adjacent tissue breakdown probably play some role in recurrence.

In the absence of specific symptoms or circumstances necessitating surgery, it is not clear that correction is indicated. One patient not operated on has been followed for more than 20 years with only mild dyspnea on exertion. A recent study of coronary fistulas¹¹ including one secondary to trauma (Case 1) concluded that there is

little anatomic or functional change associated with uncorrected lesions with small or moderate sized shunts after 17 year follow up, but that large shunts probably warrant closure.

Cardiac catheterization generally confirmed the diagnosis in the cases reported although as shown by our case the O₂ stepup may be small and insignificant. Selective coronary angiography is clearly the procedure of choice for diagnosis and delineation of these lesions.

In summary a case of traumatic coronary artery fistula is presented. Ten other cases reported in the literature were reviewed and the remarkable homogeneity of this entity was noted. The pertinent management problems including pre and post operative course, presentation, complications, indications for correction and follow up were discussed.

REFERENCES

- 1 Parmley L F, Mattingly T W and Manion W C. Penetrating wounds of the heart and aorta. *Circulation* 17:1953 1958.
- 2 Jones R C and Jahnke E J. Coronary artery atrioventricular fistula and ventricular septal defect due to penetrating wound of the heart. *Circulation* 32:1995 1965.
- 3 Tsagaris R J and Bustamante R A. Coronary arteriovenous fistula and myocardial infarction due to trauma. *Am J Cardiol* 18:177 1966.
- 4 Storey C F and Kuzman W J. Traumatic coronary artery-right atrial fistula. *Ann Thorac Surg* 4:352 1967.
- 5 Kahn D. Myocardial contusion due to steering wheel injury. *JAMA* 200:155 1967.
- 6 Sarot I A, Schechter D C and Weber D J. Post traumatic coronary arteriovenous fistula with surgical cure. *Ann Thorac Surg* 6:112 1968.
- 7 Forker A J and Morgan J R. Coronary artery fistula due to non penetrating chest trauma. *JAMA* 215:289 1971.
- 8 Lynn R B and Fay J E. Post traumatic cardiocoronary fistula. *Can J Surg* 14:335 1971.
- 9 North R I, Blake H A and Nelson W P. Coronary artery-right atrial fistula secondary to bullet wound of the heart: report of a case with successful surgical repair. *Milit Med* 136:267 1971.
- 10 Bravo A J, Glancy D I, Epstein S E et al. Traumatic coronary arteriovenous fistula: A 20 year follow up with serial hemodynamic and angiographic studies. *Am J Cardiol* 27:673 1971.
- 11 Morgan J R, Forker A B, O'Sullivan M J and Fosburg H G. Coronary arterial fistulas

seven cases with unusual features. *Am J Cardiol* 3:432 1972

Jaffee R B, Glancy L, Epstein S E, Brown B G and Morrow A G. Coronary arterial right heart fistulae. Long term observations in seven patients. *Circulation* 47:133 1973

Constantino T, Falcone M W, Perloff J K, DeLeon A C Jr and Jamison W L. Trau-

- matic coronary arterial right ventricular fistula. Report of a case. *Med Ann D C* 41:70 1972
- 14 Cheng T O and Adkins P C. Traumatic aneurysm of left anterior descending coronary artery with fistulous opening into left ventricle and left ventricular aneurysm after stab wound of chest. Report of case with successful surgical repair. *Am J Cardiol* 31:384 1973

Clinical pathologic conference

Jane Somerville MD, FRCP*

S U Khalig, MRCPath

A C Brewer FRCS

Donald Heath MD FRCP, FRCPath

Liverpool England

PROFESSOR HEATH A retired window cleaner aged 65 years was seen in Surgical Outpatients on September 14th 1972, with a history of anorexia and epigastric pain of one year's duration. The epigastric pain was not related to eating and at times lasted all day. He had lost weight for six weeks and had recently started vomiting. He had been seen at the hospital in March 1967 with a reducible right inguinal hernia which had been surgically repaired in May 1967 but which had recurred. On examination in the Outpatients Department no abnormal physical signs were found but a barium meal was carried out on September 21st. He was subsequently admitted to hospital on September 25th. No abnormal physical signs were elicited. In particular no abnormalities were found in the chest; the apex beat being felt in the fifth left intercostal space in the mid clavicular line. No murmurs were heard. The systemic blood pressure was 110/60 mm Hg and the radial pulse was 84 per minute. An operation was performed on September 27.

Three days after the operation he had a sudden attack of pain in the left hand. On examination the left hand and forearm were blue and cold and their power was reduced. There was loss of sensation to pin prick over all the fingers and there was blunting to pin prick from the level of the transverse

palmar crease to 3 cm below the elbow. The brachial and radial pulses were not palpable. The patient was not in atrial fibrillation and the radial pulse was regular. A few hours after this episode a second operation was performed.

One day later on October 2nd the patient was walking to the toilet when he developed a sudden onset of breathlessness and sudden pain in his left loin. He looked grey and his systemic blood pressure was 90/60 mm Hg. His radial pulse was 110 per minute. There was no blood in the urine. His respiratory rate was 60 per minute. There were rhonchi in the chest and there were basal crepitations. There were no abnormal clinical signs in the abdomen. There was no sign of external hemorrhage. His legs were not swollen and the calves were not tender.

During the next three to four days he developed urinary retention for which he required catheterization. It was noticed that his left leg was becoming swollen and edematous.

On October 9th he had a further episode of breathlessness accompanied by cyanosis and sweating. On examination he had basal crepitations. Despite treatment his condition gradually deteriorated and he died on October 14th.

Investigations The levels of hemoglobin

From the Medical School University of Liverpool Liverpool England

Received for publication April 4 1973

Reprint requests to Professor Donald Heath The University of Liverpool New Medical School, Ashton Street P O

Box 147 Liverpool L69 3BX England

*Senior Lecturer in Institute of Cardiology Honorary Consultant National Heart Hospital London.

Decem 1973 Vol 86 N

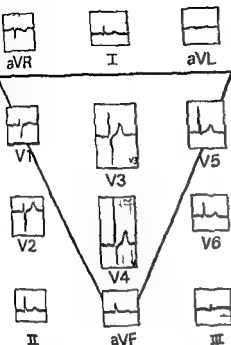


Fig. 1 Electrocardiogram of patient taken on September 26 1972



Fig. 2 Radiograph of patient's chest taken on September 26 1972

(Gm per 100 ml) packed cell volume (per cent) and mean corpuscular hemoglobin concentration (per cent) were as follows on May 11 1967 14.9 Gm % 44% 34% on September 26 1972 14.8 Gm % 44% 33% on October 10 1972 11.1 Gm % 35% 32%

The serum creatine kinase level on October 2 1972 was 760 ml U/ml (normal range in our laboratory was 0 to 130). The α -hydroxybutyric dehydrogenase level on the same day was 430 ml U/ml (normal range in our laboratory was 153 to 324). Blood group A Rh+

Urine examination on October 2 showed no evidence of hematuria albuminuria or glycosuria.

DR SOMERVILLE: I have been told that the diagnosis of the case under discussion today requires clinical acumen. If a woman is right under these circumstances it is called intuition, but if a man is right it is knowledge! Our patient had a year's history of anorexia and epigastric pain which eventually became associated with vomiting. I do not believe that his inguinal hernia is relevant because within six days of having a barium meal examination he

was on the operating table. Clearly this man had an acute severe and possibly malignant condition. I think it likely that he had a carcinoma of the stomach.

MR BREWER: That diagnosis was supported by the results of the barium meal examination which showed destruction of the normal antral mucosa and gross deformity of the prepyloric region of the stomach with some degree of stenosis.

DR KHALIQ: Later at necropsy it was confirmed that there was an ulcerating annular carcinoma 3 cm in extent in the grossly narrowed pyloric canal. Histological examination showed an adenocarcinoma of the stomach which had penetrated the full thickness of the muscle coat and metastasized to one regional lymph node.

MR BREWER: A gastroenterostomy was carried out.

DR SOMERVILLE: I note that at the time of the operation there were no abnormal signs in the chest. Furthermore the chest radiographs and electrocardiograms taken in 1967 and 1972 were normal (Figs 1 and 2). Three days after operation he had a sudden attack of pain in the left forearm and hand which were blue and cold and suffered loss of power and sensation. The brachial and radial pulses were not palpable. That adds up to only one thing: a sudden arterial occlusion. There are rela-

tively few causes of sudden occlusion of this type. He may have had a dissecting aneurysm, but in my experience this would almost certainly be associated with central chest pain if it involved the left subclavian artery and it would not present just as pain in the hand. He is more likely to have had an embolus, and presumably the second operation reported was an embolectomy. Since the brachial and radial pulses were not palpable the embolus must have been above that level and presumably was in the axillary artery.

Such emboli can arise from various sites. They may originate from thrombus in the left atrium or left atrial appendage. Thrombosis arising there is usually associated with a change of rhythm, usually atrial fibrillation. This did not apply to the patient under discussion although it is possible that he may have had a short paroxysm of fibrillation before he was seen by a doctor or nurse. Sometimes patients with mitral stenosis do throw off emboli into the systemic arteries without atrial fibrillation, but I am inclined to think that in a man of 65 years with mitral stenosis atrial fibrillation would have been present. Another source of embolism in the left atrium is a myxoma but of course in this case the surgeons would have detected fragments of myxoma and not thrombus from the brachial artery.

MR RAWSTHORNE: At the second operation the brachial artery was approached medially and the material removed was recent thromboembolus. It did not resemble the jelly like appearance of myxoma.

DR SOMERVILLE: However you would agree that fragments of myxoma can have recent thrombus superimposed on them disguising their true nature. Another source of embolus is mural thrombus following infarction of the left ventricle, but there is no evidence of that and I would discard that diagnosis. I notice the elevated enzymes but I presume this is related to the death of muscle following arterial embolism in the arm. Thrombus too can arise over atheromatous plaques on the aorta.

DR ROBERTSON: Would you expect to find a myxoma in the left atrium or trivial mitral stenosis in a man of 65 years with a heart of normal size?

DR SOMERVILLE: Yes, I believe it is possible to have a small myxoma which does not cause physical signs and recent thrombosis on such a small mass might lead to systemic embolism. We might be helped in this diagnosis by finding a raised sedimentation rate or abnormalities in the plasma proteins. With regard to mitral valve disease, I do believe one can find trivial mitral stenosis in the elderly not leading to cardiac enlargement.

PROFESSOR HEATH: Perhaps at this point I could summarize the argument so far by saying that we are seeking a source of systemic emboli occurring in a man with no clinical, electrocardiographic or radiological evidence of cardiopulmonary disease.

DR SOMERVILLE: That is right. This is a practical problem that the Cardiologist comes across from time to time. Another possibility at the back of my mind is the situation where one may have a small atrial septal defect with a paradoxical embolus from the venous site. However, I do not favor this diagnosis in view of the normal chest radiograph and electrocardiogram. It seems more likely to me that we are dealing with an embolus which has arisen from the left ventricle or systemic arteries.

Two days after the embolectomy the patient developed breathlessness, an increase in respiratory rate and a fall in systemic blood pressure. It seems likely that he had a pulmonary embolus. I notice that at this stage they found no evidence of thrombosis in the deep veins in the leg, but of course this does not necessarily exclude a pulmonary embolus because the first may occur without such clinical signs in the leg. It is also conceivable that he had a hemorrhage. The hemoglobin level fell during this period, but of course this is confused by the fact that he had a recent operation. The pain he experienced in the left loin could also have been produced by a renal or splenic infarct. As there was no hematuria I favor the latter diagnosis.

PROFESSOR HEATH: Perhaps you would like to look at the next slide which shows a picture of the man's legs as they appeared in the terminal phase of his illness.

DR SOMERVILLE: Were arterial pulses present in the left leg?



Fig 3 View of the heart from its superior aspect. The roof of the atria and the upper portion of the interatrial septum have been removed. A large thromboembolus (E) is impacted in an atrial septal defect which lies immediately beneath the rim of alveolar septum indicated by the arrow. While most of the thromboembolus (E) lies in the right atrium (RA), a small part lies in the left atrium (LA). (Original magnification $\times 13$)

MR RAWSTHORNE Yes the pulses were full

DR SOMERVILLE Well in that case I believe the cause of the swelling in that leg was a deep venous thrombosis rather than an arterial embolism

PROFESSOR HEATH Well in that case how do you fit together all the changes you have been describing? You consider that we are dealing with a deep vein thrombosis in the left leg associated with pulmonary embolism. There is systemic arterial embolism in the left arm and you describe the possibility of a splenic infarction. How do you relate all these things together?

DR HENDERSON Do you think that all these signs and symptoms described could be explained on the basis of thrombophlebitis migrans?

DR HARLEY Do you think it likely that the source of the emboli was thrombosis in the pulmonary veins?

DR SOMERVILLE I have seen tumor in pulmonary veins following extension from a bronchial carcinoma which has given rise to systemic embolism but of course we have no evidence of a tumor of lung in this case

MR POPE You say that there could not have been an atrial septal defect because it would have shown up on the radiograph or electrocardiogram but would this apply to very small defects?

DR SOMERVILLE At this age the most likely type of atrial septal defect that would allow an embolus to pass from systemic veins into systemic arteries would be a defect of secundum type but by this age of 65 years I would have anticipated that the heart would be enlarged. Furthermore I would have anticipated some evidence of right ventricular hypertrophy in the electrocardiogram. Of course one does have to bear in mind that an embolus might pass through a small valvar foramen ovale that had never given rise to symptoms or which did not reveal itself on clinical radiographic or electrocardiographic examination. However this would have been unlikely to occur unless there was some increase in pressure in the right atrium and there does not appear to be any basis for this.

MR SHATWELL Surely such a rise in pressure might have been produced by the



Fig 4 View of the heart from the left atrium (LA) after removal of the main portion of thromboembolus. The atrial septum has been cut along the line indicated by the black arrows. An atrial septal defect (white arrow) communicates with the right atrium (RA). Thromboembolus (E) is still impacted in a persistent valvar foramen ovale. (Original magnification $\times 2.5$)

thromboembolism to the lung which you consider occurred?

DR SOMERVILLE: Yes, it could. However, the emboli to the brachial artery occurred before evidence for the possible pulmonary embolism.

DR KAY: Surely pulmonary emboli can be silent?

DR SOMERVILLE: I agree, but since only five days elapsed between the time of operation and the impaction of the embolus in the left arm, one would have to postulate a lot of silent pulmonary emboli in that time to raise the pressure in the right side of the heart.

PROFESSOR HFATH: I think at this stage we could ask Dr Khaliq what he found at necropsy.

DR KHALIQ: The body was that of an elderly man of medium build and short stature (5 ft 5 in.). Partially healed gastroenterostomy and embolectomy wounds were present. The left lower limb was swollen and edematous due to thrombosis of the left femoral vein. There was bilateral

pulmonary thromboembolism. Emboli of various ages were found occluding lobar and segmental branches of both main pulmonary arteries. On histological examination, some of these emboli were partially organized. Some had blocked muscular pulmonary arteries and hence appeared to have been capable of elevating pulmonary vascular resistance. The heart was dilated and weighed 466 Gm (with attached fat). There was an elongated thromboembolus in the right atrium which was 6 cm in length and 2.5 cm in diameter. It was impacted in a small atrial septal defect and protruded into the left atrium (Fig 3). The atrial septal defect was 1 cm in diameter, anterior to the fossa ovalis and had a crescentic upper border with no septal tissue at the lower border (Fig 4). A smaller embolus (0.4 cm in diameter) was impacted in a persistent foramen ovale. Both coronary arteries were patent and showed minimal non-occlusive atheroma. The myocardium was normal. The mitral, tricuspid, aortic, and pulmonary valves were normal.

The right ventricle was very dilated

The liver was congested and sections showed centrilobular venous congestion. The spleen had a wedge shaped infarct at its lateral margin. The urinary bladder was hypertrophied due to prostatic enlargement due to benign fibroleiomyoepithelial hyperplasia.

This was a case of paradoxical embolism in which emboli arising from thrombi in the left femoral vein reached the peripheral systemic arterial circulation through defects in the atrial septum. Repeated thromboemboli also reached the lung elevating pulmonary vascular resistance and the pressure in the right atrium. This would have favored the passage of thromboemboli through the atrial septal defect and the persistent foramen ovale. The defects in the atrial septum themselves became occluded by impaction of thromboemboli. The passage of emboli through these defects or fragmentation of the impacted emboli would account for the episodes of systemic arterial embolism occluding the left brachial artery and leading to infarction of the spleen.

PROFESSOR HEATH As a pathologist one has to say that paradoxical embolism is one of those classical conditions that one includes in undergraduate teaching and yet rarely sees. It is described in most text books of pathology but how common is this condition?

DR SOMERVILLE I think it is seen very rarely in clinical practice but one does have to say that it occurs in the very type of case we have been discussing today where there is a small atrial septal defect. I have previously seen two cases of paradoxical embolism both occurring in elderly patients with deep vein thrombosis and small atrial septal defects. I think this is fascinating and certainly the pathologist has been able to explain all the signs and symptoms here on the basis of one pathology as he so frequently can but of course he has rather a selected referral of patients! However I still think that the first event in this patient was a systemic arterial em-

bolism. At that stage there was no evidence that he had had previous pulmonary embolism.

PROFESSOR HEATH Rare as this condition is it would appear to be even rarer to actually catch an embolus in the process of impaction and passage through the foramen ovale. What about the incidence of paradoxical embolism in congenital heart disease such as a ventricular septal defect or a patent ductus arteriosus?

DR SOMERVILLE This is not likely to occur because the flow of blood through such defects is in the wrong direction for it to happen namely from left to right. In the presence of pulmonary hypertension with reversal of flow it could occur. In an atrial septal defect there is part of the cardiac cycle where it is quite easy for reversal of flow of blood to take place. During atrial diastole the pressure in the right atrium may be slightly higher than in the left thus allowing a right to left shunt and then one could of course have a paradoxical embolus. I am interested in the description of the atrial septal defect. Small ostium primum defects are not necessarily associated with the characteristic electrocardiographic changes. However I would have expected some abnormality in the anterior cusp of the mitral valve.

PROFESSOR HAY I don't remember seeing cases of paradoxical embolism in congenital heart disease. How often do you get an atrial septal defect of that size with a normal electrocardiogram?

DR SOMERVILLE Well I think there is no doubt that unsuspected patency of the atrial septum is found not infrequently at necropsy. It is clear from this meeting today that such patency can lead very rarely to paradoxical embolism. If this man had not had the carcinoma and associated deep vein thrombosis it seems very unlikely that his small atrial septal defect would have caused him any trouble at all or perhaps even come to light.

DIAGNOSIS Paradoxical embolism
 Atrial septal defect
 Carcinoma of stomach

Fundamentals of clinical cardiology

Straight back syndrome

Clinical and hemodynamic study of 9 cases

Shuzo Matsuo, MD

Masato Ioshioaka MD

Katsusuke Yano MD

Kunitake Hashiba, MD

Nagasaki Japan

Rawlings¹ first described the straight back syndrome as a cause of pseudo heart disease in 1960 and additional reports have followed.²⁻⁴ We have studied 9 cases of straight back syndrome finding a small systolic pressure gradient between the right and main pulmonary arteries in one catheterized case. The purpose of this paper is to describe the clinical and hemodynamic findings of these 9 cases and to discuss in detail the one having the pulmonary arterial pressure gradient.

Material and methods

Nine patients (4 males and 5 females) with straight back syndrome were studied between January 1964, and April 1968 in the First Department of Internal Medicine, Nagasaki University School of Medicine Japan. Ages ranged from 12 to 18 years (average 15.6 years). Two patients were referred with the diagnosis of atrial septal defect (Cases 5 and 8). The others were referred for evaluation of cardiac murmurs. The criteria used for patient selection were a straight thoracic spine on inspection and a lateral chest x ray. A com-

plete history and physical examination as well as electrocardiograms (ECG's) and phonocardiograms (PCG's) were obtained. In each patient the ratio of the anteroposterior dimension to the transthoracic dimension was calculated from the chest x ray as described by De Leon and associates.⁵ Right heart catheterization was performed in 6 patients including right sided angiocardiology in 4.

Results

The physical chest roentgenographic ECG and PCG findings are summarized in Table I. All patients had absence of the normal kyphotic curvature of the thoracic spine on inspection (Fig. 1). Two patients also had mild pectus excavatum (Cases 5 and 8, Table I). On physical examination all patients had a systolic ejection murmur Grade 2-3, 6 in intensity at the pulmonary area. Splitting of the second sound was observed in all cases varying from 0.03 to 0.06 sec on expiration and increasing with inspiration. There were no other notable physical findings. PCC recorded in the second intercostal space at the left sternal

From The Third Department of Internal Medicine Nagasaki University School of Medicine Nagasaki Japan
Received for publication March 2, 1969

Reprint requests to Dr. Shuzo Matsuo, The Third Department of Internal Medicine Nagasaki University School of Medicine 7-1 Sakamoto Machi Nagasaki Japan

Table 1 The physical chest roentgenographic PCG and ECG findings in 9 patients with straight back syndrome

| Patient no | Initial age and sex | Straight back | Pectus excavatum | Chest x-ray | | | | PCG | | ECG | | |
|------------|---------------------|---------------|------------------|------------------|------------------|--------------------|-------------------------|----------|-------------------------|-----|-----------------------|---------------------|
| | | | | Prominence of PA | Left sided shift | Pancake appearance | AP/trans thoracic ratio | Syst mur | Split of S ₂ | RAD | rSr in I ₁ | Clock wise rotation |
| 1 | Y O 12 F | + | - | + | - | + | 30% | + | + | + | - | - |
| 2 | M T 13 M | + | - | - | - | + | 35% | + | + | - | + | - |
| 3 | N K 14 F | + | - | + | - | + | 35% | + | + | + | + | + |
| 4 | H N 16 F | + | - | + | + | + | 33% | + | + | - | - | - |
| 5 | S S 10 F | + | + | + | - | - | 35% | + | + | - | - | - |
| 6 | M K 17 F | + | - | + | - | - | 35% | + | + | + | - | - |
| 7 | H K 17 M | + | - | + | - | - | 35% | + | + | - | - | + |
| 8 | T I 17 M | + | + | + | - | - | 37% | + | + | - | + | + |
| 9 | Y T 18 M | + | - | + | - | - | 35% | + | + | + | - | - |

Abbreviations M = male F = female PA = pulmonary artery AP/trans thoracic ratio = anteroposterior diameter ratio PCG = phonocardiogram Syst mur = systolic murmur Split of S₂ = splitting of the second sound ECG = electrocardiogram RAD = right axis deviation.

border are shown in Fig 2. The ECG findings were an axis varying from 90 to 110 degrees in 4 patients and an rSr pattern (r ≥ r) in I₁ and I₂ in two other patients (Fig 3). One additional patient had both of these findings. Chest x-ray revealed prominence of the pulmonary artery conus in 8 patients (Fig 4) and left sided shift of the heart shadow in one patient (Fig 5). There was a pancake appearance of the heart in 4 patients (Figs 4 and 5). The ratio of the anteroposterior diameter to the transthoracic diameter varied from 27 per cent to 37 per cent. These ratios are decreased below normal in all except one case (Case 6).

A summary of the right heart catheterization data in 6 patients including angiocardiology in 4 is shown in Table II. A slight increase in right ventricular systolic pressure (> 30 mm Hg) was observed in 3 patients. Five patients had a small resting systolic pressure gradient (4 to 13 mm Hg) across the right ventricular outflow tract. No cases showed arteriovenous shunting. Angiocardiology was performed in 4 patients. Three of these showed flattening of the right ventricular outflow tract (Fig 6) and two showed compression of the anterior wall of the right ventricle against the sternum (Fig 7).



Fig 1 View of the posterior thorax in Case 3. The dorsal spine lacks the normal outward curve.

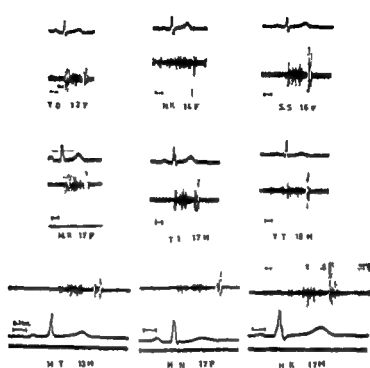


Fig 2 Phonocardiogram of all 9 cases. All are recorded at the left sternal border in the second intercostal space on expiration

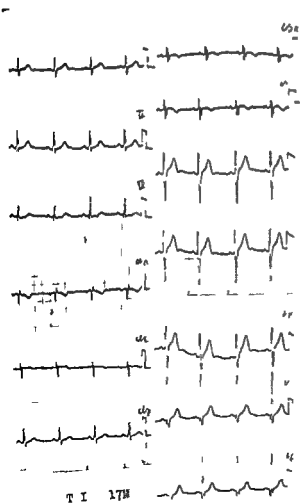


Fig 3 Electrocardiogram in Case 8

In Case 1 although there was no clear stenosis of the main right pulmonary artery on angiocardiography an abrupt systolic pressure gradient of 10 mm Hg was observed in the main right pulmonary artery as the catheter was withdrawn from the peripheral right pulmonary artery to the main pulmonary artery (point A to point B Fig 7). This pressure change was recorded where the right pulmonary artery passed anteriorly to the straight thoracic spine. This maneuver was performed three times (Fig 8) with identical results.

Discussion

Straight back syndrome is sometimes misdiagnosed as organic heart disease most frequently atrial septal defect, idiopathic pulmonary artery dilatation or mild pulmonary stenosis.^{1,2,5} Two of our cases (Cases 5 and 8) were thought to have atrial septal defect because of the findings of a wide splitting of the second sound and an ejection systolic murmur in the pulmonic area, prominence of the pulmonary conus on the chest x ray film, and an rSr' pattern in Leads V_1 and V_{2R} of the ECG.

All but one of our cases showed prominence of the pulmonary artery segment on chest x ray. This finding as well as the leftward shift and pancake appearance of the



Fig 4 Anteroposterior chest x ray in Case 1 showing prominence of the pulmonary artery. Lateral view shows a straight dorsal spine and narrow anteroposterior diameter with distinct retrocardiac space.



Fig 5 Anteroposterior view of the chest x ray in Case 2 showing leftward displacement of the heart and slight prominence of the pulmonary artery. Lateral view shows a straight dorsal spine and narrow anteroposterior diameter.

heart may be explained by rotation and displacement due to narrow anteroposterior diameter of the thorax in this syndrome. In all but one of our cases the ratios of the anteroposterior to transverse diameters of the chest x ray were under 37 per cent significantly less than the values described as normal by De Leon and associates.⁴ These investigators suggested that the systolic murmur in the second left intercostal space may relate both to displacement of the main pulmonary artery

and to an exaggeration of the normal ejection vibration due to the proximity of the vessel to the chest wall.

In this regard angiocardigraphic study showed compression of the right ventricle and proximity of the pulmonary artery to the chest wall in two cases (Fig 7) and flattening of the outflow tract of the right ventricle in three cases (Fig 6).

Association of pectus excavatum with the straight back syndrome was reported in one of twelve cases by Rawlings² and two

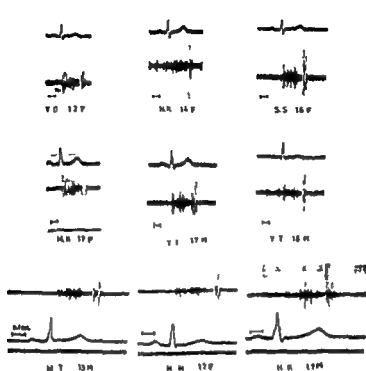


Fig 2 Phonocardiograms of all 9 cases. All are recorded at the left sternal border in the second intercostal space on expiration.

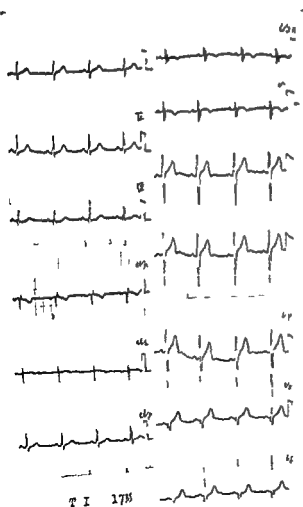


Fig 3 Electrocardiogram in Case 8

In Case 1, although there was no clear stenosis of the main right pulmonary artery on angiocardiography, an abrupt systolic pressure gradient of 10 mm Hg was observed in the main right pulmonary artery as the catheter was withdrawn from the peripheral right pulmonary artery to the main pulmonary artery (point A to point B, Fig 7). This pressure change was recorded where the right pulmonary artery passed anteriorly to the straight thoracic spine. This maneuver was performed three times (Fig 6) with identical results.

Discussion

Straight back syndrome is sometimes misdiagnosed as organic heart disease, most frequently atrial septal defect, idiopathic pulmonary artery dilatation, or mild pulmonary stenosis.^{1,2,5} Two of our cases (Cases 5 and 6) were thought to have atrial septal defect because of the findings of a wide splitting of the second sound and an ejection systolic murmur in the pulmonary area, prominence of the pulmonary conus on the chest x-ray film, and an rS₁ pattern in Leads V₁ and V_{2R} of the ECG.

All but one of our cases showed prominence of the pulmonary artery segment on chest x-ray. This finding as well as the leftward shift and pancake appearance of the



Fig 7 Anteroposterior (left) and lateral view (right) of the right ventricular angiogram in Case 1. The anterior wall of the right ventricle is compressed against the sternum in the lateral view.

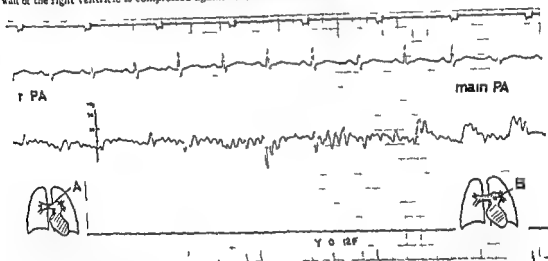


Fig 8 Continuous pressure tracing as the catheter was withdrawn from point A in the right pulmonary artery to point B in the main pulmonary artery (Case 1). A and B correspond to the same points on the angiogram in Fig 7.

catheter tip did not appear to be wedged on fluoroscopy and the difference in pressure occurred in the main branch of the right pulmonary artery when the catheter was continuously withdrawn from a peripheral pulmonary artery. Small systolic pressure gradients between the pulmonary branches and main pulmonary artery may be functional in conditions with large pulmonary blood flow due to left to right shunt.¹¹ However this patient had no arteriovenous shunt by blood oxygen studies or on angiocardiography. Although mild

congenital pulmonary artery stenosis^{12, 13} cannot be excluded as a cause of the pressure gradient we believe that compression of the right main pulmonary artery as it crosses the straight thoracic spine better explains the gradient in this case. Leinbach and co-workers⁸ have reported one case of straight back syndrome with pulmonary venous obstruction. They concluded that this was due to sternospinal cardiac compression.

This brief report also serves to re-emphasize a clinically recognizable entity which



Fig 6 Anteroposterior (left) and lateral view (right) of the right ventricular angiogram in Case 8. The flattening of the right ventricular outflow tract is seen in the lateral view.

Table II Right heart catheterization data in 6 patients, including angiocardiology in 4 patients

| Patient initials age, and sex | RA (mm Hg) | | | RV (mm Hg) | P1 (mm Hg) | L/R shunt | ACG | | |
|----------------------------------|------------|---|------|---------------|----------------------|--------------|---------------------|----------------------|------------------------------|
| | a | v | mean | | | | PA valv stenosis | Flat of RV outlet | Comp of ant wall of RV |
| Y O 12 F | 7 | 7 | 4 | 28/0-7 | mPA 24/8 rPA 14/7 | - | - | + | + |
| N K, 14 F | 5 | 5 | 4 | 30/0-5 | 26/10 | - | - | ± | - |
| S S 16 F | 7 | 5 | 4 | 30/0-7 | 30/10 | - | - | + | + |
| M K 17 F | 6 | 5 | 4 | 35/0-4 | 28/10 | - | - | - | - |
| T I 17 M | 5 | 4 | 3 | 38/0-4 | 25/10 | - | - | + | - |
| Y T 18, M | 5 | 5 | 3 | 38/0-5 | 25/8 | - | - | - | - |

Abbreviations: RA = right atrium; RV = right ventricle; PA = pulmonary artery; L/R shunt = left to right shunt; mPA = main pulmonary artery; rPA = right pulmonary artery; PA valv stenosis = pulmonary valvular stenosis; Flat of RV outlet = flattening of the right ventricular outflow tract; Comp of ant wall of RV = compression of the anterior wall of the right ventricle against the sternum; ACG = angiocardiology.

of 23 cases by Gooch and colleagues.⁷ In the present study, two of nine cases had associated mild excavatum.

In Case 1, although there was no clear narrowing of the right pulmonary artery on the angiocardiology, a mild abrupt systolic pressure gradient was observed on pressure recording when the catheter was pulled back from the peripheral right pulmonary artery to the main pulmonary artery. This gradient occurred at the point where the right main pulmonary artery crossed in front of the straight thoracic

spine (Fig 8). To our knowledge there has been no previous report of such a finding in association with this syndrome.

Brum and associates⁹ found false pressure gradients in the pulmonary arteries could result from (1) wedging of the catheter tip in a small pulmonary artery branch, (2) partial obstruction of the flow of blood by the catheter in the artery catheterized, and (3) the small, though normal differences in pressure that may be found in various regions of an anatomically normal pulmonary artery system. In the present case the

Appraisal and reappraisal of cardiac therapy

Edited by Arthur E. DeGross and Julian Frieden

Therapeutic uses of atrial pacing

Seymour Furman MD
Bronx N Y

Ventricular cardiac pacing has a secure well-established and well-defined place in the treatment of heart disease and cardiac arrhythmias. Atrial pacing to the contrary is irregularly used and with a slowly evolving group of indications. The reasons for the difference are:

1 The difficulty in establishing consistent atrial pacing and sensing¹

2 The paucity of atrial or ventricular arrhythmias controllable by atrial pacing relative to those amenable to ventricular pacing and the realization that the effects of atrial arrhythmia usually can be ameliorated or adequately treated by the more readily applied stable and reliable ventricular pacing

3 The presence almost everywhere in the heart of a ventricular signal at least as large and frequently much larger than the atrial signal so that pacemaker triggering from the atrial signal alone may be erratic and sensing from either the atrial or the ventricular signal or both may occur². Atrial activity can fail to recycle an atrial pacemaker or it can be recycled by a QRS complex effectively slowing the rate of atrial stimulation

4 The existence of intermittent or partial atrioventricular block so that the ventricle may not at all times be affected by atrial

pacing sometimes when the circumstances most require³

5 The late development of atrial fibrillation or flutter which negates the effect of atrial pacing on the ventricle

Nevertheless indications for atrial pacing exist in which stimulation of the atria is an excellent alternative to ventricular pacing⁴ and others for which atrial pacing is to be preferred to ventricular pacing⁵

The major therapeutic indications for atrial pacing are:

I Bradycardia caused by an inadequate sinus or atrial rate in the presence of intact atrioventricular conduction

A Episodic sinus bradycardia sinus arrest or sinoatrial block with asystole

B As above with periods of supra-ventricular tachycardia—i.e. the bradytachy syndrome

II Suppression of ventricular irritability by an overriding conducted atrial rhythm

III Atrioventricular synchrony to increase cardiac output

A By direct atrial pacing in the presence of intact A-V conduction

B By atrioventricular sequential

From the Cardiac Service, Division of Surgery, Mount Sinai Hospital and Medical Center, Bronx, N.Y.
Supported in part by United States Public Health Service Grant HE-04666-13

Received for publication April 12, 1973

Reprint requests to Seymour Furman, MD, Cardiac Service, Division of Surgery, Mount Sinai Hospital and Medical Center, E. 61st St. & 1st Ave., Box 11, New York, N.Y. 10022

should be borne in mind when certain mild congenital heart lesions are suspected

Summary

Clinical, electrocardiographic, radiologic, and hemodynamic findings in 9 cases of straight back syndrome are described. In one case a small systolic pressure gradient was observed as the catheter was pulled back from the peripheral right pulmonary artery to the main pulmonary artery. To our knowledge this is the first report of a gradient associated with this syndrome. The possible hemodynamic causes are discussed. It is re-emphasized that the cardiac findings in straight back syndrome frequently mimic those of certain mild congenital heart defects.

The authors are grateful to Professor Yoshito Takaoka, The First Department of Internal Medicine, Nagasaki University School of Medicine, and to Dr Aaron B. Shiffer and Dr Howard C. Cohen, Heart Station, Michael Reese Hospital, Chicago, Ill., for reviewing the manuscript.

REFERENCES

- 1 Rawlings M S The straight back syndrome: a new cause of pseudo-heart disease *Am J Cardiol* 5:333 1960
- 2 Rawlings M S Straight back syndrome: a new heart disease *Dis Chest* 39:435 1961
- 3 Serratto M and Kezdi P Absence of the physiologic dorsal kyphosis: Cardiac signs and hemodynamic manifestations *Ann Intern Med* 58:938 1963
- 4 Datey K K, Deshmukh M M, Engineer S D, and Dilvi C P Straight back syndrome *Br Heart J* 26:614 1964
- 5 De Leon A C Jr, Perloff J K, Twigg H, and Wajid M The straight back syndrome: Clinical cardiovascular manifestations *Circulation* 32:193 1965
- 6 Rubenstein H J and Johnson R B The effect of external compression on the murmur and thrill of the straight back syndrome *Am Heart J* 74:88 1967
- 7 Gooch A S, Maranhao V, and Goldberg H The straight thoracic spine in cardiac diagnosis *Am Heart J* 74:595 1967
- 8 Leinbach R C, Harthorne J W, and Dinsmore R E Straight back syndrome with pulmonary venous obstruction *Am J Cardiol* 21:588 1968
- 9 Baum D, Khoury G H, Ongley P A, Swan H J C, and Kincaid O W Congenital stenosis of the pulmonary artery branches *Circulation* 29:680 1964
- 10 Zimmerman H A *Intravascular catheterization* 2nd ed. Springfield, Ill. 1966. Charles C. Thomas Publishers.
- 11 Shaffer H A and Bliss H A Pulmonary artery stenosis *Am J Med* 26:517 1959
- 12 Luan L L, D'Silva J L, Casal H M, and Dillon R F Stenosis of the right main pulmonary artery: clinical, angiocardigraphic and catheterization findings in ten patients *Circulation* 21:1116 1960
- 13 Agustsson M H, Arrillaga R, Gavil B M, Bricoff J P, Nassif S I, and Lendrum B L The diagnosis of bilateral stenosis of the primary pulmonary artery branches on characteristic pulmonary trunk pressure curves: A hemodynamic and angiocardigraphic study *Circulation* 26:421 1962
- 14 Schlesinger F G and Meester G T Supra-valvar stenosis of the pulmonary artery *Br Heart J* 29:829 1967
- 15 Rios J C, Walsh B J, Matsumi R A, Sims A J, and Ewy G A Congenital pulmonary artery branch stenosis *Am J Cardiol* 24:1518 1969

Appraisal and reappraisal of cardiac therapy

Edited by Arthur C DeGraff and Julian Frieden

Therapeutic uses of atrial pacing

Seymour Furman MD
Bronx NY

Ventricular cardiac pacing has a secure well-established and well-delineated place in the treatment of heart disease and cardiac arrhythmias. Atrial pacing to the contrary is irregularly used and with a slowly evolving group of indications. The reasons for the difference are:

1 The difficulty in establishing consistent atrial pacing and sensing¹

2 The paucity of atrial or ventricular arrhythmias controllable by atrial pacing relative to those amenable to ventricular pacing and the realization that the effects of atrial arrhythmia usually can be ameliorated or adequately treated by the more readily applied stable and reliable ventricular pacing

3 The presence almost everywhere in the heart of a ventricular signal at least as large and frequently much larger than the atrial signal so that pacemaker triggering from the atrial signal alone may be erratic and sensing from either the atrial or the ventricular signal or both may occur.² Atrial activity can fail to recycle an atrial pacemaker or it can be recycled by a QRS complex effectively slowing the rate of atrial stimulation

4 The existence of intermittent or partial atrioventricular block so that the ventricle may not at all times be affected by atrial

pacing sometimes when the circumstances most require³

5 The late development of atrial fibrillation or flutter which negates the effect of atrial pacing on the ventricle

Nevertheless indications for atrial pacing exist in which stimulation of the atria is an excellent alternative to ventricular pacing⁴ and others for which atrial pacing is to be preferred to ventricular pacing⁵

The major therapeutic indications for atrial pacing are:

- I Bradycardia caused by an inadequate sinus or atrial rate in the presence of intact atrioventricular conduction
 - A Episodic sinus bradycardia sinus arrest or sinoatrial block with asystole
 - B As above with periods of supra-ventricular tachycardia—i.e. the bradytachy syndrome
- II Suppression of ventricular irritability by an overriding conducted atrial rhythm
- III Atrioventricular synchrony to increase cardiac output
 - A By direct atrial pacing in the presence of intact A-V conduction
 - B By atrioventricular sequential

From the Cardiothoracic Service Division of Surgery, Montefiore Hospital and Medical Center, Bronx, NY.
Supported in part by United States Public Health Service Grant HE 04666-13.
Received for publication April 12, 1973.

Reprint requests to Seymour Furman, MD, Cardiothoracic Service Division of Surgery, Montefiore Hospital and Medical Center, E. G. Hall & Balbridge Avenue, Bronx, NY 10467.

pacing in the presence of A V block

IV Rapid atrial pacing to terminate supraventricular and re entry tachycardias

Physiology of atrial pacing

Atrioventricular synchrony usually produces greater cardiac output than ventricular paced rhythms.⁶ In the presence of complete heart block, atrioventricular asynchrony with lack of the atrial contribution reduces output compared to A V synchrony.⁷ If conduction is present (ventricular pacing for sinus bradycardia) then retrograde stimulation of the atrium and atrial contraction during ventricular systole against a closed atrioventricular valve reduces cardiac output even further.⁸

In the normal, however, the contribution of atrial systole during right atrial pacing increases cardiac output little compared to direct ventricular pacing.⁹ Exercise in the normal during atrial or ventricular pacing at similar rates causes similar increases in stroke volume and cardiac output.¹⁰

In those with heart disease unlike the normal, increasing cardiac rates with atrioventricular synchrony usually produces increases in cardiac index, time tension index, ventricular power, mean systolic pressure and systolic ejection compared to ventricular pacing at similar cardiac rates.^{11, 12} Exercise in those with cardiac disease produces greater increments in cardiac output with atrial compared to ventricular pacing at similar rates.¹³

Atrial pacing requires less ventricular work to produce more rapid ejection time, greater left ventricular end diastolic pressure and aortic systolic pressure than ventricular pacing.¹⁴ Atrial synchronous pacing and in all likelihood atrioventricular sequential pacing produce cardiac outputs 15 to 20 per cent greater than continuous direct ventricular pacing under these circumstances.¹⁵

After cardiac surgery and acute myocardial infarction maximum cardiac efficiency is required and atrioventricular synchrony at rapid (80 to 100) physiologic rates can produce cardiac outputs up to 60 per cent greater than ventricular pacing

at the same rate. In the presence of insufficiency of the A V valves, especially the mitral, direct ventricular pacing may decrease cardiac output. Stroke volume and arterial pressure for a single beat decrease when atrium and ventricle are out of phase and increase when fortuitous synchrony occurs. In the presence of A V valvular insufficiency during ventricular pacing the asynchronous ventricular contractions will be associated with reductions of output either for a single or for a succession of several asynchronous beats.¹⁶ Reductions of stroke volume may be to 50 per cent of normal levels. If block is frequent the total cardiac output drop may be profound. If it is infrequent the patient may be almost as symptomatic during periods of apparently normal cardiac pacing as before the pacer was inserted.¹⁷

This event referred to as "pacemaker syndrome" is most effectively managed by the restoration of A V synchrony.¹⁸ In the presence of normal atrial rate and heart block, atrial synchronous (atrial triggered ventricular pacing) and in the presence of sinus bradycardia or arrest, atrioventricular sequential pacing are both satisfactory solutions to the problem.¹⁹ If sinus bradycardia without heart block exists then direct atrial pacing is satisfactory.²⁰ Direct corrective action toward mitral or tricuspid valve may be required.

The atrial electrode

Implantation of an atrial electrode by thoracotomy is not difficult and several approaches exist. The flat spiral atrial epicardial electrode used as an atrial sensor for atrial synchronous pacing can be used to sense and pace the atrium. Electrodes designed for transvenous ventricular stimulation can be adapted for suture to and stimulation of the atrium.²¹

The problem of transvenous atrial electrode placement has not been definitively resolved though occasional papers with small successful series appear. Ventricular stimulating electrodes may be placed in the coronary sinus from which stable atrial pacing and intermittently satisfactory sensing is possible.²² A J shaped electrode may be placed into the atrial appendage and is also effective for pacing, though less so for

sensing the P wave^{26, 27} An electrode with wires projecting (harpoon like) from its tip may be introduced within a larger plastic catheter with a lumen into the atrial appendage²⁸ The lumen catheter is large and difficult to manipulate and the author has failed to place it in about half the attempts An ingeniously designed electrode introduced from the femoral vein via a modified Ross needle perforates and grasps the atrium septum²⁷ This approach leaves the connection to a pacemaker in the lower abdomen as the electrode is brought subcutaneously through the groin Connection to a ventricular electrode as well as to the atrial seems difficult should atrial synchronous or A V sequential pacing be desired

Modes of atrial pacing

I *Asynchronous atrial pacing* Stimulation of the atrium is continuous the pace maker does not respond to atrial or ventricular activity The rhythm is useful as it avoids the problems of sensing spontaneous atrial activity and can best be used when fixed sinus bradycardia with satisfactory A V conduction at least at physiologic rates exists Such pacing can be applied via any of the electrode placement techniques

II *Atrial triggered pacing* Two major subgroups exist These are atrial triggered ventricular pacing commonly referred to as atrial synchronous or P wave triggered pacing which will not be discussed further and atrial triggered (non competitive) atrial pacing

In atrial triggered atrial pacing atrial activity modifies the pacemaker activity in a fashion analogous to the modification of ventricular pacing by ventricular activity

A *ATRIAL INHIBITED ATRIAL PACING* Stimulation of the atrium occurs if the atrial rate is below the pacemaker escape rate Increase in the spontaneous atrial rate inhibits pacemaker stimulation A relatively conventional pulse generator may be used though its sensitivity should be to a one millivolt signal rather than to the more conventional two millivolt (ventricular) signal

Two problems may arise (1) the in-

creased pulse generator sensitivity may still be inadequate to sense atrial activity and pacemaker competition with atrial activity may occur yielding pacemaker induced premature atrial contractions and (2) the ventricular signal as indicated earlier is large throughout the heart As most inhibited pulse generators have short refractory periods (the longest for widely used generators is 200 msec) the possibility exists that following atrial stimulation the pacer may be inhibited and be recycled by the subsequent QRS complex

B *ATRIAL SYNCHRONIZED ATRIAL PACING* In this instance a synchronous pulse generator is employed It too should be of one millivolt sensitivity and the same electrode sensing problems exist as for an inhibited pacer However as synchronous pacemakers have a refractory period of 350 to 400 msec it is unlikely that the QRS complex will occur early enough to recycle the pulse generator An additional advantage is that transcutaneous increase of pulse generator rate is readily possible and that atrial rates up to 140 per minute can be imposed²⁹ if required to override a tachycardia

III *Atrioventricular sequential* When sinus activity is sufficiently slow that increase in atrial rate is desirable but A V conduction is compromised or unreliable pacing is required for both atrium and ventricle especially when the atrial contribution to cardiac output is of great consequence³⁰ Four circumstances exist

1 Acute myocardial infarction with heart block and sinus bradycardia these patients obviously require the maximum of cardiac efficiency and cardiac output can be increased by the restoration of atrioventricular synchrony compared to direct ventricular pacing³⁰

2 Postoperative cardiac surgery the same circumstances exist as following myocardial infarction³¹

3 With more profound degrees of myocardial failure the atrial contribution plays a greater role in cardiac output In severe congestive heart failure a physiologically timed atrial contraction will provide a much greater cardiac output than will asynchronous ventricular pacing (see above)³²

4 With severe insufficiency of the A V

valves especially the mitral direct ventricular pacing accentuates the degree of insufficiency and cardiac output may fall rather than rise with an increase in cardiac rate. If the atrial rate is adequate, atrial synchronous pacing will ameliorate the problem, if not A V sequential pacing is required.

A V sequential pacing may be at a fixed atrial rate, a fixed A V delay and with asynchronous ventricular pacing assuming that no competitive spontaneous ventricular activity will occur. A more satisfactory approach especially when the possibility exists either of spontaneous atrial activity with conduction or premature ventricular activity is such pacing with ventricular inhibition in approach called *Bifocal demand** (atrioventricular sequential ventricular inhibited) pacing.²²

The Bifocal demand pacer designed both as an external and as an implantable model has at least in its implantable form suffered from serious technical problems which have severely restricted its utility.

IV Rapid atrial pacing To this point pacing of the atrium has assumed a physiologic rate of stimulation analogous to ventricular pacing at a similar rate. A unique approach to atrial pacing is that of rapid stimulation of the atrium at rates of 400 to 1 200 per minute with no attempt to achieve A V conduction.²³ The ventricle is protected from such rates by the inability of the conduction system to transmit such rapid impulses. The attempt is to convert a pathologic atrial rhythm to regular sinus rhythm.²⁴ Rapid atrial pacing, as asynchronous manually applied and terminated Rapid atrial pacing operates in one of three distinct fashions. In the presence of an atrial or junctional tachycardia the pacemaker rate may be set above the atrial rate capture and then allow slowing. Alternatively a continuous and stable atrial arrhythmia is converted to an unstable rhythm which reverts to regular sinus rhythm and finally circus or reciprocating pathways may be interrupted by the rapid pacemaker stimuli.

1 Atrial flutter Atrial flutter can frequently be converted to atrial fibrillation

or regular sinus rhythm by rapid atrial pacing. Wide variability of results exists: one report was of success in 48 of 68 attempts with 21 converted to sinus rhythm and 27 to atrial fibrillation of which sixteen ultimately returned to sinus rhythm.²⁵ In another group of 15 patients seven resumed atrial flutter with the cessation of atrial pacing and six remained in atrial fibrillation.²⁶ In the remaining two capture of the atrium never occurred. In all direct current cardioversion was eventually required. Rapid atrial pacing is a secondary procedure for atrial flutter but may be attempted for its ease of application without the use of anesthesia and in the presence of digitalis intoxication without the danger of ventricular tachy fibrillation.²⁷

2 Supraventricular tachycardia Supraventricular tachycardia both atrial and nodal may not be controlled by overriding atrial pacing within the physiologic range (100 to 180 per minute).²⁸ Even pacing at a rate in excess of the atrial rate may not capture control and break the tachycardia. Rapid atrial pacing converts the tachycardia to atrial fibrillation an unstable rhythm which then often reverts to a regular rhythm. Both discrete impulses and 60 Hz alternating current have been used successfully with similar results.^{29, 30}

Increases of atrial rate by rapid atrial pacing may not produce atrial fibrillation but in atrial tachycardia with increasing atrioventricular block—i.e. 3:1 to 5:1 It may be necessary to administer digitalis and propranolol in conjunction with pacing in order to achieve this effect. A permanently implanted radio frequency pacemaker operated as necessary by the patient may be extremely useful for recurrent tachycardias.³¹

3 Re entry tachycardia and pre excitation syndrome In both instances pacing of either the atrium or the ventricle at a near physiologic rate may not capture the rate or even if it does may not allow slowing or cessation of pacing without return to the original rhythm. Rapid atrial pacing may convert the rhythm to unstable atrial fibrillation or may interrupt the circus movement.³² As the approach is not universally effective its efficacy in a specific instance should be demonstrated prior to

establishing permanent pacing in the case of a recurrent tachyarrhythmia such as Wolff Parkinson White syndrome⁴⁴ Some patients will require the addition of small to moderate doses of suppressive drugs such as beta adrenergic blocking agents to facilitate cardiac sensitivity

In the Wolff Parkinson White syndrome it is probable that the reciprocating pathways or circus movements responsible for the tachycardia may be interrupted by the rapid stimulation While reports of termination of the tachycardia by a single atrial stimulus have appeared⁴⁵ a period of several seconds of rapid atrial pacing is more effective at a practical therapeutic level

⁴ Ventricular arrhythmias and especially tachycardias have been successfully managed by ventricular pacing at rates adequate to overdrive and suppress the arrhythmia⁴⁶ In the presence of intact conduction atrial overdrive is probably more effective

a The electrode may contribute to further ventricular irritability by mechanical stimulation of the ventricle This is avoided by atrial stimulation

b Cardiac output is increased by atrial relative to ventricular pacing and this increase in output may assist in resolution of the etiology of the tachycardia Additional drug or other therapy should be directed at the heart as overdrive suppression of tachycardia is a short term expedient

Conclusion

In an area more limited than that for ventricular pacing atrial pacing at physiologic rates or at rapid rates for short periods of time has multiple significant uses The major limitations of atrial pacing for long term use are the lack of availability of reliable atrial leads and electrodes which will function well as atrial sensors as well as stimulators Because of the simultaneous existence of atrial arrhythmia and atrioventricular block an A-V sequential pacer of proved reliability will undoubtedly increase the utility and use of atrial pacing

REFERENCES

- 1 Furman S Reicher-Reiss H and Escher D J W Atrio-ventricular sequential pacing and pacemakers Chest 63:183 1973

- 2 Nathan D A Center S Wu C Y et al An implantable synchronous pacemaker for the long term correction of complete heart block Circulation 27 682 1963
- 3 Lemberg L Castellanos A and Berkovits B Pacing on demand in AV block JAMA 191 12 1965
- 4 Silverman L F Mankin H T and McGoon D C Surgical treatment of an inadequate sinus mechanism by implantation of a right atrial pacemaker electrode J Thorac Cardiovasc Surg 50 264 1968
- 5 Moss A J Rivers R J Jr and Griffith L S C Transvenous left atrial pacing for recurrent ventricular fibrillation N Engl J Med 2 8 928 1968
- 6 Kosowsky B D Scherlag B J and Damato A N Reevaluation of the atrial contribution to ventricular function study using His bundle pacing Am J Cardiol 21 518 1968
- 7 Samet P Bernstein W H Nathan D A et al Atrial contribution to cardiac output in complete heart block Am J Cardiol 16 1 1965
- 8 Furman S and Escher D J W Principles and techniques of cardiac pacing New York 1970 Harper & Row Publishers 165
- 9 Benhimol A Elli J G and Dimond E G Hemodynamic consequences of atrial and ventricular pacing in patients with normal and abnormal hearts Am J Med 39 911 1965
- 10 Stein E Damato A N and Kosowsky B D Cardiovascular responses to alterations in heart rate above and below the sinus rate Am J Cardiol 17 140 1966
- 11 Martin R H Cobb L A Lau S H et al Impaired cardiac function during ventricular pacing in man Clin Res 13:122 1965
- 12 Samet P Castullo C and Bernstein W H Hemodynamic consequences of atrial and ventricular pacing in subjects with normal hearts Am J Cardiol 18 522 1966
- 13 Samet P Castullo C Bernstein W H et al Hemodynamic results of right atrial pacing in cardiac subjects Dis Chest 53 133 1968
- 14 Gilmore J P Sarnoff J Mitchell J H et al Synchronicity of ventricular contractions observations comparing hemodynamic effects of atrial and ventricular pacing Br Heart J 23 799 1963
- 15 Benhimol A and Liggett M L Cardiac hemodynamics during stimulation of the right atrium right ventricle and left ventricle in normal and abnormal hearts Circulation 33 933 1966
- 16 Chamberlain D A Leimbach R C Vassaux C E et al Sequential atrioventricular pacing in heart block complicating acute myocardial infarction N Engl J Med 282 577 1970
- 17 Samet P Hemodynamic sequelae of cardiac arrhythmias Circulation 47 399 1973
- 18 Mitani T Mizuno A Hasegawa T et al Atrial rate as an indicator for optimal pacing rate and the pacemaker syndrome Ann Cardiol Angiol 10 371 1971
- 19 Smyth N P D Bacos J M Massumi R A

- et al Permanent transvenous synchronous cardiac pacing *Chest* 59:493 1971
- 20 Moss A J Rivers R J and Cooper M Long term percutaneous atrial pacing from the proximal portion of the coronary vein *JAMA* 209 543 1969
- 21 Nussbaum M and Levit S Technique for permanent implantation of atrial pacemaker *Surgery* 68:916 1970
- 22 Perryman R A and Seely W C Permanent atrial pacing *Ann Thorac Surg* 15:16 1973
- 23 Kramer D H and Moss A J Permanent percutaneous atrial pacing from the coronary vein *Circulation* 42:427 1970
- 24 Smyth N P D Keshishian J M Brusa A P et al Permanent transvenous atrial pacing *Ann Thorac Surg* 11:360 1971
- 25 Zucker I R Parsonnet V and Gilbert L A method of permanent transvenous implantation of an atrial electrode *Am Heart J* 85:195 1973
- 26 Porstmann W Witte J Dressler L et al P wave synchronous pacing using anchored atrial electrode implanted without thoracotomy *Am J Cardiol* 30:74 1972
- 27 Udall J A Cardiac pacing by means of an attached endocardial electrode a case report *Curr Ther Res* 13 660 1971
- 28 Furman S Escher D J W Parker B et al Electronic analysis for pacemaker failure *Ann Thorac Surg* 8:57 1969
- 29 Benchimol A and Goldstein M R A review of atrial pacing clinical and laboratory applications *Ann N Y Acad Sci* 167:604 1969
- 30 Leinbach R C Chamberlain D A Kistor J A et al A comparison of the hemodynamic effects of ventricular and sequential A V pacing in patients with heart block *Am Heart J* 78 502 1969
- 31 Woodson R D Atrial pacing in cardiac surgical management *Northwest Med* 66 1109 1967
- 32 Samet P Bernstein W H and Levine S Significance of the atrial contribution to ventricular filling *Am J Cardiol* 15:195 1965
- 33 Castillo A Berkovits H V Castellanos A Jr et al Bifocal demand pacing *Chest* 59 360 1971
- 34 Haft J I Kosowsky B D Lau S H et al Termination of atrial flutter by rapid electrical pacing of the atrium *Am J Cardiol* 20:239 1967
- 35 Haft J I Lau S H Stein E et al Atrial fibrillation produced by atrial stimulation *Circulation* 37:70 1968
- 36 Vergara G S Hildner F J Schoenfeld C H et al Conversion of supraventricular tachycardias with rapid atrial stimulation *Circulation* 46:788 1972
- 37 Rosen K M Sinno M Z Gunnar R N et al Failure of rapid atrial pacing in the conversion of atrial flutter *Am J Cardiol* 29 524 1972
- 38 Kistor J A DeSanctis R W and Leinbach R C Long term percutaneous atrial pacing *Circulation* 40:535 1969
- 39 DeSanctis R W Diagnostic and therapeutic uses of atrial pacing *Circulation* 43:748 1971
- 40 Lister J W Cohen L S Hildner F J et al Electrical stimulation of the atria in patients with an intact atrioventricular conduction system *Ann N Y Acad Sci* 167:785 1969
- 41 Lister J W Gulotta S J Keller J W et al Treatment of supraventricular tachycardia by right atrial alternating current stimulation *Am J Cardiol* 29:208 1972
- 42 Davidson R M Wallace A G Sealy W C et al Electrically induced atrial tachycardia with block *Circulation* 44:1014 1971
- 43 Barold S S Linhart J W Samet P et al Supraventricular tachycardia initiated and terminated by a single electrical stimulus *Am J Cardiol* 24:37 1969
- 44 Lau S H Stein E Kosowsky B D et al Atrial pacing and atrioventricular conduction in anomalous atrioventricular excitation (Wolff Parkinson White Syndrome) *Am J Cardiol* 19 354 1967
- 45 Durrer D Schaal L and Schuilenburg R M Role of premature beats in the initiation and termination of supraventricular tachycardia in the Wolff Parkinson White syndrome *Circulation* 36 644 1967
- 46 Sowton E Leatham A and Carson P The suppression of arrhythmias by artificial pacemaking *Lancet* 2:1098 1964

Annotations

Tricyclic antidepressants and cardiac disease

Cardiovascular complications following the administration of tricyclic antidepressants are well documented.¹ The most serious of these deaths was reported in association with imipramine by Kristiansen in 1961 and with amitriptyline by the Australian Drug Evaluation Committee in 1963. Suspicion that the tricyclic antidepressants were cardiotoxic was increased by the number of serious adverse cardiac effects reported to the Committee on Safety of Drugs² by 1968 and epidemiological evidence to confirm or refute this suspicion was sought.

A hospital based drug information system described by Coull and associates³ which links details of patient identification with diagnosis and drug therapy made possible the rapid identification of all patients in the Aberdeen General Hospital who had received amitriptyline in the forty month period ending September 1971. The number of patients identified was 864, this being the total population at risk of the adverse effect. Since however Kristiansen⁴ had shown that patients with cardiac disease were more susceptible to cardiac side effects detailed study was limited to patients who had cardiac disease and who had received amitriptyline in the two weeks preceding discharge from hospital or death, as it seemed unlikely that earlier discontinuation of the tricyclic antidepressant drug could be associated with subsequent death. Therefore 119 patients were the subject of further study and of these 23 patients had died. Examination of the clinical records revealed that 13 of the deaths were sudden and unexpected—the death occurred within 24 hours of the onset of the terminal illness, there being no previous indication that death was imminent.

To ascertain whether amitriptyline played a role in inducing these deaths it is necessary to compare them with the deaths in a control population who did not receive amitriptyline. Since there is a very close correlation between sex, diagnosis and mortality rate it is of paramount importance when looking for an event such as unexpected death that the patients should be adequately matched for these factors. In addition as the chance of recording a death in hospital is related to the length of hospital stay this parameter was also included in the matching criteria. It proved possible because of the large numbers of patients in the system (120,000 patients September 1971) to find 339 controls of the same sex, very similar age and diagnoses and period of hos-

pitalization but who did not receive a tricyclic antidepressant. Fifteen patients died in this group, 3 of them unexpectedly, as compared with 13 in the amitriptyline group and this difference is possibly significant ($\chi^2 = 1.5$, $p < 0.05$, single tail). The occurrence of myocardial infarction, cardiac failure and severe anemia (Hb < 70 per cent) were used to estimate the severity of the patients' cardiac disease and general condition in both groups. It was similar and the controls were therefore also well matched in these respects.

The drugs administered to the amitriptyline population and their controls were carefully reviewed. While a large number of different drugs was prescribed for each group and those who died in each group received on average more than those who lived the overall prescribing pattern was similar. Amitriptyline was prescribed in the test group in conventional dose regimes and digoxin was the only drug in common use in both groups which had a direct cardiac effect.

The progress and treatment of the amitriptyline group and their controls was followed after discharge from hospital. A proportion of the amitriptyline patients continued to receive the drug and it was of interest to note that the drug had been prescribed for one of the controls reinforcing the assumed adequacy of the control population. The ultimate prognosis in terms of mortality rate in the original test and control groups with or without amitriptyline subsequently proved to be remarkably similar and suggests that amitriptyline may have precipitated by a number of months or years an inevitable sudden death.

The occurrence of sudden unexpected death was also investigated in 88 patients without cardiac disease who received amitriptyline and a carefully matched control population. One sudden unexpected death attributed to a cerebrovascular accident occurred in the amitriptyline group and none in the controls. Imipramine is related pharmacologically to amitriptyline and it therefore seemed possible that imipramine might also be cardiotoxic. Four sudden unexpected deaths occurred in 87 cardiac patients receiving imipramine compared with 2 in a control group. These numbers are too small to permit valid conclusions but it seems possible that imipramine may be associated with unwanted cardiac effects and this is now the subject of further study.

The investigation of major adverse drug reactions such as death poses a difficult problem for it is not easy to justify prospective controlled clinical trials of potentially lethal drugs. In this situation the comparison of retrospective data from a hospital drug information system for two groups of patients adequately matched but for the administration of the suspected drug has been valuable and the suspicion that amitriptyline may be associated with sudden unexpected death has been confirmed in a population with pre-existing cardiac disease. While such epidemiological investigations can contribute to the safer and more rational use of drugs, they can never establish with certainty a cause and effect relationship and detailed pharmacological investigation is necessary. Nevertheless even if the pharmacological basis is not fully understood in view of the serious nature of the adverse effect in question, the results suggest that amitriptyline should be prescribed with caution in patients with cardiac disease and that similar pharmacological agents such as imipramine merit further investigation.

D C Nair MD
Lecturer in Social Medicine
University of Aberdeen
Foresterhill
Aberdeen Scotland

REFERENCES

- 1 Meyler L. Side effects of drugs V. Amsterdam 1966. Excerpta Medica Foundation.
- 2 Smith R B and Rusbatch B J. Amitriptyline and heart block. *Br Med J* 3:311 1967.
- 3 Edwards A L. Imipramine myocardial toxicity. *N Y State J Med* 2:1979 1964.
- 4 Kristiansen E S. Cardiac complications during treatment with imipramine (tofranil). *Acta Psychiatr Neurol Scand* 36:127 1961.
- 5 Australian Drug Evaluation Committee. Adverse reactions to amitriptyline. *Med J Aust* 1:741 1965.
- 6 Inman H H W. Personal communication 1968.
- 7 Coull D C, Crooks J, Davidson J F, Gallon S C and Weir R D. A method of monitoring drugs for adverse reactions I. Methyl dopa and haemolytic anemia. *Eur J Clin Pharmacol* 3:46 1970.

Hemoglobin, plasma lipids, and coronary heart disease

During recent years several so-called risk factors for coronary heart disease (CHD) have been identified—and intensively discussed. Many seem to have used the word risk factor to denote not only a definite risk in the individual patient but also a direct pathogenic influence of the factor itself. Clearly a factor the presence of which is associated with increased rate of development of CHD may on one hand cause this development but might on the other only be a parallel or associated phenomenon without direct pathogenic importance. In spite of the fact that a large number of risk factors for CHD have been identified, little is known about the way in which they are associated with the development of CHD. Much more research is needed to clarify these matters. One approach is to study the possible interrelationship between risk factors.

The first large prospective study to identify risk factors was the Framingham study,¹ which has been followed by similar investigation aimed at identifying risk factors. One of the more recent ones is the Stockholm Prospective Study,^{2,3} which has confirmed previous findings of elevated serum cholesterol, elevated blood pressure (systolic as well as diastolic), elevated hemoglobin values, and smoking as risk factors for CHD and has added two others, elevated fasting serum triglycerides and erythrocyte sedimentation rate. The linear relationship found

between plasma cholesterol and triglycerides respectively and the rate of development of new events of CHD is shown in Figs 1 and 2.

High hemoglobin or hematocrit (packed cell volume) has been identified as a risk factor for CHD.^{4,5} As it has not been a matter of extreme values and as it is known—i.e. from polycythemic conditions—that the rheological properties of the blood are not influenced until the hemoglobin values reach very high levels (above 20 Gm per 100 ml), it has been difficult to understand the meaning of these findings. A recently reported study⁶ might throw some light on the problem.

As already mentioned in the Stockholm Prospective Study, plasma lipids (cholesterol and triglycerides) were identified as risk factors for CHD but so were high hemoglobin values.

A statistical analysis of a possible relationship between plasma lipids and hemoglobin in the Stockholm Prospective Study has now been performed. It was shown that there exists a weak but significant ($P < 0.01$) positive correlation between on one hand hemoglobin and on the other plasma cholesterol or plasma triglycerides (Table 1). This correlation has previously been found for cholesterol in anemic women but not in healthy women with normal hemoglobin values and not in healthy men. No such correlation has earlier been found between

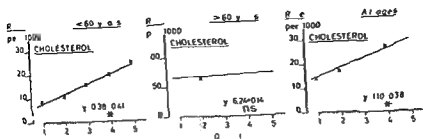


Fig. 1 Linear regression analysis of the rate of new events of CHD in relation to plasma cholesterol quintiles

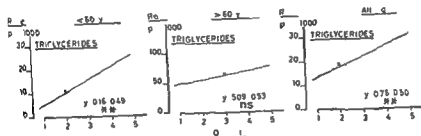


Fig. 2 Linear regression analysis of the rate of new events of CHD in relation to plasma triglyceride quintiles

plasma triglycerides and hemoglobin. Also partial correlation coefficients, keeping age constant, clearly show that the correlation exists regardless of age.

It is of interest that in our material the differences in cholesterol values corresponding to variations in hemoglobin within the normal female range (11 to 14 Gm per 100 ml) were of the same order of magnitude as the variations found by Elwood and associates⁷ in his material of anemic women (or 6 mg per 100 ml for each gram of hemoglobin per 100 ml blood) (Table I).

The variations in serum lipid values with the hemoglobin levels may of course be coincidental and largely independent of each other. They could however also be explained by changes in plasma volume. If we assume that in a woman the blood volume remains unchanged as the hemoglobin decreases from 14 Gm to 11 Gm per 100 ml the packed red cell volume would decrease from 40 to 32 per cent and consequently the plasma volume would increase from 60 to 68 per cent. If the total plasma pool of lipids would remain unchanged the concentration of cholesterol would diminish from an average 250 to 220 mg per 100 ml and of triglyceride from 110 to 97 mmole/L, that is a decrease with 30 mg per 100 ml and 0.13 mmole/L respectively. From the regression equations (Table I) the corresponding figures can be calculated to be 18 mg/100 ml for cholesterol and 0.13 mmole per liter for triglycerides. Similar calculations with similar results may be carried out for men. Although many assumptions are involved in such calculations—and great standard errors occur in the equations—simple changes in plasma volume might at least in part account for the positive correlation between hemoglobin and plasma lipid. It should be emphasized that this is a theory or speculation today without any proof. There are of course also other factors such as an increased synthesis of

Table I Regression equation between hemoglobin values (x) and serum lipid levels (y) in men and women (aged 20 to 54 years)

| Variables | Regression equation |
|--------------------------|----------------------|
| Men (n = 1 190) | |
| Hemoglobin-triglycerides | $y = 0.124x - 0.262$ |
| Hemoglobin-cholesterol | $y = 4.75x + 208.0$ |
| Women (n = 895) | |
| Hemoglobin-triglycerides | $y = 0.044x + 0.605$ |
| Hemoglobin-cholesterol | $y = 6.00x + 198.5$ |

porphyrins and lipoproteins as recently suggested by Lees and colleagues.⁸

However irrespective of why hemoglobin is positively correlated to both plasma cholesterol and triglycerides the correlation may in part explain why a high hemoglobin value has been found to be a risk factor for CHD. The difference in mean hemoglobin values between the lowest and highest quintile in the Stockholm Prospective Study was approximately 4.0 Gm per 100 ml for men. The regression equations in Table I then predict that the cholesterol and triglyceride levels (on this average) would be $4.0 \cdot 4.75 = 19$ mg per 100 ml and $4.0 \cdot 0.124 = 0.50$ mmole per liter higher in the top than in the bottom hemoglobin quintile.

The rate of CHD had increased from 30 per 1 000 in the lowest to 44 per 1 000 in the highest quintile. For men 50 to 70 years of age the average part of quintiles 2, 3 and 4 was for cholesterol 31 mg per 100 ml and for triglycerides 0.34 mmole per liter

The regression equations (Figs 1 and 2) thus predict—in this region—that the rate of new events of CHD will increase with 0.12 per 1 000 and 0.15 per 1,000 for an increase of 1 mg per 100 ml of cholesterol and 0.01 mmole per liter of triglycerides respectively. Applying these figures to the above mentioned values of 19 mg per 100 ml and 0.50 mmole per liter they would correspond to an increase in the rate of CHD events of 2.3 and 7.5 respectively or together 9.8 per 1 000. This value arrived at from calculations using regression equations with statistical as well as biological limitations is fairly close to the actual difference of 14 per 1 000 between the bottom and top hemoglobin quintile. Thus hemoglobin is a risk factor for CHD might be explained not by the high hemoglobin value itself but by the fact that the plasma lipids are increased as well.

It should finally also be stressed that so called risk factors for CHD to a large extent covariate and are difficult to treat as single phenomena. As an example we can mention the erythrocyte sedimentation rate (ESR) which was found to be a risk factor. It has long been known that ESR and hemoglobin values are closely related (for discussion see Ref. 8). Furthermore recent studies have indicated that the ESR is closely associated also with other risk factors. Thus studies on the Stockholm Prospective Study material have shown that there exist positive correlations between elevated ESR values and elevated plasma lipids (cholesterol as well as triglycerides¹⁹) and that elevated ESR values are found in persons with asymptomatic hyperlipidemia.²⁰ The final answer to the question of the importance of elevated hemoglobin or erythrocyte sedimentation values as pathogenetic or non pathogenetic risk factors for CHD must await the results of further studies.

L. E. Bottiger MD
Professor of Medicine
Department of Medicine
Karolinska Hospital
Stockholm, Sweden
L. A. Carlson MD
Professor and Head
Department of Geriatrics
University of Uppsala
Uppsala, Sweden

REFERENCES

- 1 Kannel W B, Dawber R R, Kagan A, Rebovsky N and Stokes J. Factors of risk in the development of coronary heart disease—six year follow up experience. *Ann Int Med* 53:33 1961.
- 2 Carlson L A and Lindstedt S. The Stockholm Prospective Study I. The initial values for plasma lipids. *Acta Med Scand Suppl* 493:1968.
- 3 Bottiger L E and Carlson L A. The Stockholm Prospective Study. New events of coronary heart disease in men in relation to findings at initial examination. 9 year follow up. Stockholm 1972. Skandia International Symposium (In press).
- 4 Carlson L A and Bottiger L E. Ischaemic heart disease in relation to fasting values of plasma triglycerides and cholesterol. *Lancet* 1:865 1972.
- 5 Dawber R R and Kannel W B. Susceptibility to coronary disease. *Mod Concepts Cardiovasc Dis* 30:671 1961.
- 6 Bottiger L E and Carlson L A. Relation between serum cholesterol and triglyceride concentrations and haemoglobin values in non anaemic healthy persons. *Br Med J* 3:731 1972.
- 7 Elwood P C, Mahler R, Sweetman P, Moore F and Welsby E. Association between circulating haemoglobin level, serum cholesterol and blood pressure. *Lancet* 1:589 1970.
- 8 Bottiger L E and Svedberg C A. Normal erythrocyte sedimentation rate and age. *Br Med J* 2:85 1967.
- 9 Lees R S, Song C S, Ievers R D and Kappas A. Hyper betalipoproteinemia in acute intermittent porphyria. *N Engl J Med* 282:432 1970.
- 10 Bottiger L E. Erythrocyte sedimentation rate and plasma lipids. *Acta Med Scand* 193:53 1973.
- 11 Bottiger L E, Carlson L A and Olsson A G. Elevation of erythrocyte sedimentation rate in asymptomatic hyperlipidemia. *Br Med J* (In press 1973).

Aspirin-like drugs and prostaglandins

The aspirin like drugs are a diverse group of compounds all of which possess in some measure the antipyretic, anti-inflammatory and analgesic actions which are characteristic of aspirin.

Aspirin itself was introduced into clinical medicine in the late 1890's but decoctions of the bark or leaves of willow trees which contain derivatives of the parent compound (salicylic acid) have been valued for their medicinal properties for centuries.

The leaves are bitter, reads one account, and the Hottentots and the Arabs use them in rheumatic fever.¹ Yet despite their long history and wide spread use (it is estimated that 4 000 million aspirin tablets are consumed each year in England alone) the question of how the aspirin like drugs exerted their therapeutic effect remained unanswered. Although they were known to interfere with a wide variety of cellular enzymes, no convincing relation

hip could be established between the experimental findings and the observed clinical actions. This situation changed in 1971 when a group of workers at the Royal College of Surgeons of England led by J. R. Vane discovered that aspirin like drugs inhibited the biosynthesis of prostaglandins.¹⁻⁴ Prostaglandins are a family of long chain fatty acids which possess an unusually wide spectrum of biological activity. The capacity for generating these extremely potent substances is apparently shared by all tissues of the body and at the time of this discovery there was some evidence suggesting that prostaglandins participated in the pathogenesis of fever and inflammation. Thus as Vane pointed out inhibition of prostaglandin synthesis might explain the mode of action of the aspirin like drugs. Today almost two years later all the available evidence supports this conclusion.^{5,6}

Almost all the aspirin like drugs have been found to inhibit the enzyme system which synthesizes prostaglandins (prostaglandin synthetase). The steroidal anti-inflammatory drugs and the narcotic analgesics which have some similar therapeutic actions are ineffective as are many other drugs which do not have any aspirin like actions and this suggests that inhibition of prostaglandin synthesis is a property peculiar to this class of drugs. In man a single dose of salicylates or indomethacin is sufficient to reduce the synthesis of prostaglandin by an easily detectable amount and treatment with therapeutic doses of these drugs for two or three days results in a very pronounced inhibition.

There are several lines of evidence that point to the involvement of prostaglandins in inflammation. Prostaglandins are found in inflammatory exudate and if injected intradermally reproduce many of the salient features of the inflammatory response. Furthermore the delayed phase of inflammation which is selectively suppressed by aspirin is the phase in which prostaglandins appear. The relative therapeutic activity of several well known anti-inflammatory drugs fits well with their anti PG synthetase activity and the concentrations required for inhibition are within the plasma levels achieved by these drugs during treatment. With regard to the antipyretic activity of these drugs (a central action) prostaglandins have been detected in the CSF of rats during fever and when this was reduced by the administration of aspirin like drugs there was also a concomitant decrease in the concentration of prostaglandin. Conversely when injected into the CSF of animals or systematically into man prostaglandins cause a fever which is not blocked by aspirin like drugs.

The foregoing findings are consistent with the notion that the prostaglandins are causal agents in inflammation and fever and thus inhibition of synthesis could adequately account for two of the three therapeutic effects of these compounds—but what of their analgesic activity? At first this was more difficult to accommodate within the theory. These drugs possess only a weak analgesic action and are only effective against certain types of low intensity pain. It was known that prostaglandins were painful when injected into man and some animals but only in much larger concentrations than would be likely to occur in tissues. However when contin-

uously infused subdermally in doses too small to cause overt pain the prostaglandin caused a long lasting hyperalgesia—an increased sensitivity to mechanical stimulation as well as to other endogenous substances such as histamine or bradykinin. It seems from these studies that prostaglandins released away from the site of a local lesion could sensitize afferent pain fibers to other mechanical or chemical stimuli. Thus the analgesic action of the aspirin like drugs could be attributed to the removal of this facilitating effect and this would explain why aspirin is only a weak analgesic.

The aspirin like drugs have a number of unexpected side-effects such as gastric irritation, nephrotoxicity and disturbances in collagen metabolism and there are already several lines of evidence which link these effects with their anti PG synthetase activity. For example prostaglandins could be released from the gastric mucosa during the churning of the stomach contents after a meal and since some of the prostaglandins are known to be inhibitors of gastric acid secretion a negative feedback loop has been postulated which controls the pH of the gastric contents. If this loop were broken by the administration of prostaglandin synthesis inhibitors acid secretion could increase leading perhaps to irritation or erosion of the gastric mucosa a side-effect often associated with these drugs.

The discovery that the aspirin like drugs inhibit prostaglandin synthesis has given the clinician an insight into the nature as well as the treatment of pain, fever and inflammation—but it has also been of value to the physiologist who wishes to assess the function of the prostaglandins in the normal body and also to the enzymologist who wishes to learn more about the mechanism of synthesis of these fatty acids.

One apparent anomaly in the hypothesis has recently been resolved with interesting consequences. Paracetamol (acetaminophen, APAP) an aspirin like drug which has analgesic and antipyretic but no anti-inflammatory actions was inactive against the prostaglandin synthetase prepared from peripheral tissues such as skin or spleen against which the standard drugs were active. Did it work by a different mechanism? The only alternative appeared to be that paracetamol exerted its actions only within the CNS. When a prostaglandin synthesizing system was prepared from brain tissue it was found that the drug was indeed very active against prostaglandin synthesis.⁷ This finding suggests that the enzymes responsible for prostaglandin biosynthesis in the various tissues of the body may be differentially sensitive to inhibition by the aspirin like drugs. It may even be possible to produce drugs of this type which have selective actions on certain organs or which are free from unwanted side-effects.

In conclusion it is pertinent to ask what relevance these findings have to cardiology. This is difficult to answer at the moment. Prostaglandins have a positive inotropic effect on the myocardium of the isolated heart and increase the cardiac output and coronary flow in man and animals. More than one study suggests that they may be involved in autoregulation of coronary blood flow.⁸ At the time of writing however no disease condition has been

specifically associated with aberrant prostaglandin production by the heart

R J Flower
Department of Pharmacology
Royal College of Surgeons of England
Lincoln's Inn Fields
London WC2A 3JN England

REFERENCES

- 1 Watt J M and Breyer Brandwyk M G The medicinal and poisonous plants of Southern Africa Edinburgh 1932 E & S Livingstone Ltd
- 2 Vane J R Inhibition of prostaglandin synthesis as a mechanism of action for aspirin like drugs *Nature New Biol* 231 732 1971
- 3 Smith J H and Wilks A L Aspirin selectively inhibits prostaglandin production in human platelets *Nature New Biol* 231 235 1971
- 4 Ferreira S H Moncada S and Vane J R Indomethacin and aspirin abolish prostaglandin release from the spleen *Nature New Biol* 231 237 1971
- 5 Vane J R Prostaglandins and the aspirin like drugs *Hosp Practice* 7 61 1972
- 6 Vane J R Prostaglandins in the inflammatory response in Inflammation mechanisms and control Ed by I H Lepow and P A Ward Academic Press Inc New York and London 1972
- 7 Flower R J and Vane J R Inhibition of prostaglandin synthesis in brain explains the anti pyretic activity of paracetamol (4 acetamidophenol) *Nature* 240 410 1972
- 8 Karim S M M and Somers K Cardiovascular and renal actions of prostaglandins in Karim S M M ed Prostaglandins progress and research Oxford and Lancaster 1972 Medical and Technical Publishing Co Ltd

Surgical versus medical management of coronary heart disease

Comparisons of the operative risks of medical and surgical management of ischemic heart disease are summarized in Table I. No surgeon will ever top the surgical risks of the cardiologists. Therefore the evaluation of the surgical treatment is reduced to the answers to two questions, namely:

1 What is the survival rate when surgery is employed?

2 Do the benefits of surgery justify the risks of suffering and costs? If so, where is the evidence?

George E Burch MD
Department of Medicine
Tulane University
School of Medicine
1430 Tulane Ave
New Orleans La 70112

Table I Comparisons of operative risks of medical and surgical management of ischemic heart disease

| | Treatment | |
|---|-----------------------|---------|
| | Surgical | Medical |
| Operative mortality | 4 to 75 + % | Zero |
| Operative complications: early and late (myocardial infarction, hemorrhage, infection, thromboembolism, etc.) | 25% | Zero |
| Total cost of operation | \$8,000 to \$35,000 + | Zero |
| Closure of shunt | 25% in 2 years | Zero |
| Cures | Zero | Zero |
| Operative pain and suffering | 100% | Zero |
| Operative psychic stress to patient and family | 100% | Zero |
| Survival time | Unknown | Known |

Letters to the Editor

Echocardiographic studies of mitral valve

To the Editor

I was very interested to read the article by Millward McLaurin and Craig¹ in a recent issue of *THE JOURNAL* (AM HEART J ■ 413 1973). I would like to make the following comments on their article.

1. Measurements of mitral valve excursion (MVE) are wrong: these were measured between points D and E on the mitral valve echogram and they should be measured between points C and E, the most posterior and the most anterior points of the anterior mitral valve leaflet movement. The authors compared their measurements to those done by Wharton and Bescos.² In Wharton and Bescos' paper (p 348) it is well documented that their measurements were done between points C and E (Fig 5 postop LCG). Most measurements of the mitral valve excursion are based today on the distance between C and E³ and not D and E as done by the authors. From the above it is understood that all of Millward McLaurin and Craig's measurements of the mitral valve excursion and the numbers they compared their measurements to were wrong since the measurements are based on wrong criteria.

2. The authors state that in 16 of 17 patients with congestive cardiomyopathy (CCM) the mitral valve echogram was abnormal in that the two mitral valve leaflets never completely approximated one another during ventricular systole. This is not documented in their echocardiograms.

2A. In Fig 2 (p 414) only the anterior mitral leaflet is demonstrated during diastole. Unless both leaflets are well demonstrated during diastole, no conclusions as to the posterior leaflet's movements during systole can be drawn. From this echogram alone it seems that there is some abnormal systolic anterior movement of the anterior mitral leaflet, documented in case with HISS but seen in some patients with the mid systolic click-late systolic murmur syndrome at the upper part of the anterior mitral leaflet near the ring.

2B. In Fig 4 (p 417) (note the description of Fig 5 belong to Fig 4 and vice versa) in the third echogram (B W) it is well demonstrated that the two mitral leaflets do merge into one line during systole—the other parallel line could most probably be given by tilting the transducer. It seems according to our experience as well as to others^{4,5} that systolic separation of the mitral leaflets without posterior displacement of one or the other leaflet is not diagnostic of mitral insufficiency due to papillary muscle dysfunction. In eight of our babies with severe CCM, six of whom had and two of whom did not have mitral regurgitation, no correlation could be found between mere systolic separation (without systolic posterior displacement of the mitral leaflet) and mitral insufficiency. In

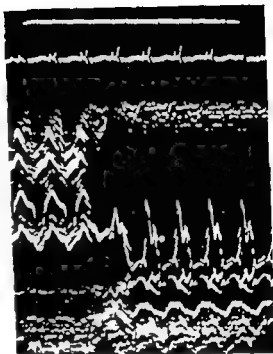


Fig 1 Echocardiogram of a 13 month old girl with severe congestive cardiomyopathy. The transducer was tilted from the aorta (left side of the picture) to the mitral valve and the left ventricle while the picture was taken. The left ventricle is very dilated, the mitral valve apparatus is dislocated posteriorly, the septum is dislocated anteriorly. The baby died and the autopsy showed a very dilated left ventricle with endocardial fibroelastosis.

fact, systolic separation of the mitral leaflets without posterior displacement can be found in many normal individuals.

3. The authors state that all of the patients with acute mitral regurgitation due to ruptured chordae tendinae showed systolic separation (p 415). In Fig 5 (p 417) case B G shows only systolic separation (which could be normal) but no posterior systolic displacement of any of the leaflets. Thus this echogram is not necessarily diagnostic of papillary muscle dysfunction.

4. One important finding which is not mentioned by the authors but is seen in two of their echograms (patients J W and B W, Fig 4) is the posterior displacement of the mitral valve apparatus in patients with CCM. In our material all eight patients with CCM demonstrated this phenomenon. The mitral valve leaflets are seen very close to the posterior wall of the dilated left ventricle away from the interventricular septum. This can be best documented by moving the transducer from the aorta to the left ventricle. The mitral apparatus instead

of being continuous with the posterior wall of the aorta is pushed posteriorly. The interventricular septum which in the normal is continuous with the anterior wall of the aorta is pushed in interior direction (Fig 1).

This echographic picture is diagnostic of CCM and demonstrates severe dilatation of the left ventricle.

Joram Glaser M.D.
Department of Pediatrics
Case Western Reserve University
2103 Adelbert Rd.
Cleveland Ohio 44106

REFERENCES

- 1 Millward D K, McLaurin I P and Craige E. Echocardiographic studies of the mitral valve in patients with congestive cardiomyopathy and mitral regurgitation. *Am Heart J* 85:413 1973.
- 2 Wharton C I P and Bescos L I. Mitral valve movement: a study using ultrasound technique. *Br Heart J* 32:344 1970.
- 3 Solinger R, Libi J and Vinhas K. Echocardiography in the normal neonate. *Circulation* 47:108 1973.
- 4 Kirber R E, Isieff D M and Hancock E W. Echocardiographic patterns in patients with the syndrome of systolic click and late systolic murmur. *New Eng J Med* 281:691 1971.
- 5 Sweetman T, Selzer A, Haimatzakis M and Cohn K. Echocardiographic diagnosis of mitral regurgitation due to ruptured chordae tendinae. *Circulation* 46:580 1972.

Reply

To the Editor

We appreciate the opportunity to comment on some of the questions raised by Dr. Glaser in his letter referring to our article published in the March 1973 issue of the *AMERICAN HEART JOURNAL*.

1. The anterior motion of the closed mitral valve during systole (C to D on the echocardiogram) is thought to be secondary to motion of the mitral annulus and does not represent independent motion of either anterior or posterior mitral valve leaflets. It seems to us that in measuring excursion of the anterior leaflet in early diastole a more appropriate convention is to measure points D to E. It is probably not justifiable to compare our results with those of Wharton and Bescos² since they made their measurements in a different way but we do agree that our measurements are wrong. The method we used is clearly stated. The technique and results are internally consistent and the relationship of early diastolic excursion of the anterior leaflet among the various patient groups is as stated in our manuscript.

2. We do not believe it to be possible to state which leaflet is prolapsing in Fig. 2. The temporal

relationship of the leaflet separation to the systolic click and subsequent murmur recorded phonocardiographically was very precise and reproducible. We would agree that the echo is not typical of the posterior bowing motion more commonly seen.¹

2B. In patient B. W. the mitral leaflet echoes do come into contact in systole but separate signals representing anterior and posterior leaflets remain apparent. It is true that multiple echo signals usually from the anterior leaflet can be produced by tilting the ultrasonic transducer so that the valve leaflet is not perpendicular to the full beam of ultrasound. In view of this potential for artifact we would agree that multiple echoes are not necessarily diagnostic of leaflet prolapse and thank Dr. Glaser for re-emphasizing this fact. Nonetheless, we were impressed and in fact stimulated to do the study by the consistency with which both mitral valve leaflet echoes could be obtained in systole in patients with CCM. Although the use of M mode scans and strip chart recorder has increased our ability to obtain good echograms of both mitral leaflets in other patients, it has remained true that a clear representation of both leaflets with multiple systolic echoes is most easily obtained in patients with CCM. Systolic separation of the leaflets is clearly not a specific sign of papillary muscle dysfunction although in our opinion this is the most tenable hypothesis to account for mitral regurgitation in CCM.

3. The first part of the echogram in patient G. A clearly shows both anterior and posterior leaflets. The upper systolic line is continuous with the posterior leaflet echo seen in diastole.

4. In most adults the posterior mitral valve leaflet is very close to the posterior wall of the LV; the anterior leaflet is somewhat variable in its relationship to the IV septum. We thank Dr. Glaser for pointing out the interesting phenomenon seen in his experience with children. We think it is pertinent to mention that a lack of continuity of the aorta and mitral apparatus or posterior dislocation of the mitral apparatus is most frequently seen in cases of double outlet RV. Although the echocardiogram of Fig. 4 shows the mitral valve leaflets to be close to the posterior surface of the heart, definite statements regarding true dislocation of the mitral apparatus require continuous scanning as pointed out by Dr. Glaser. We are currently using strip chart M mode scanning techniques to re-evaluate adult patients with dilated left ventricles in hopes of detecting changes in the geometric relationship of various cardiac structures in dilated hearts. It is our current belief that such alterations will depend more on changes in the position of the IV septum and LV posterior wall than the annular and atrioventricular valves being attached to the rather unyielding fibrous skeleton of the heart.

David A. Millward M.D.
Lambert P. McLaurin M.D.
Ernest Craige M.D.
School of Medicine
Department of Medicine
The University of North Carolina
Chapel Hill N.C. 27514

REFERENCES

- 1 Zak A, Nasser W A and Feigenbaum H Study of mitral valve action recorded by reflected ultrasound and its application in the diagnosis of mitral stenosis *Circulation* 37:789 1968
- 2 Wharton C F and Bescoe L L Mitral valve movement: a study using an ultrasound technique *Br Heart J* 32:344 1970
- 3 Kerber R E, Isaacs D M and Hancock E W Echocardiographic patterns in patients with the syndrome of systolic click and late systolic murmur *N Engl J Med* 283:691 1971

Heparin and venous thrombosis

To the Editor

I am quite disturbed by the editorial appearing in the April issue of THE JOURNAL by Dr J R O'Brien (*Am Heart J* 85:435 1973) on the question of heparin and venous thrombosis. Dr O'Brien merely repeats what he had written in the March 1973 issue of *Modern Concepts of Cardiovascular Disease*. In both instances he finds it difficult to comprehend the reason for the success of small doses of subcutaneous heparin in the prevention of venous thrombosis and embolism in the operative patient. It is a program first advocated by me in 1962 and not by Mr Kalkar who merely called it low dose heparin and altered the heparin schedule slightly. My program continues unceasingly and I still prefer to designate it as small dose heparin prophylaxis. Dr O'Brien concedes that heparin is effective according to Mr Kalkar and other English investigators who had the opportunity denied us here (FDA restriction) in evaluating their patients for deep leg vein thrombosis with the ¹²⁵I fibrinogen uptake test but according to his latelet and other studies for the wrong reasons. This assessment leads only to greater confusion in understanding the rationale and will discourage the use of heparin prophylaxis in the great need of prevention of fatal thromboembolism. However there is a rationale for the subcutaneous small dose heparin routine which prompted the program initiated by me. It is based on my earlier observation on the long overlooked phenomenon of the pulmonary megakaryocytes. This I explained fully in an earlier report to the AMERICAN HEART JOURNAL in December 1970* and other earlier publication.* But curiously enough Dr O'Brien and the British investigators Kalkar and colleagues, Gordon Smith and associates, Nicolaides and co-workers and Gallus and colleagues fail to mention this rationale. It is Mr Kalkar especially who offers no rationale and his use of low dose subcutaneous heparin. All admit that most laboratory tests are insensitive in determining such rationale as to time and dose and the effect are valueless in establishing such routine. However in my hand the modified Dale and Laidlaw Coagulometer has proved to be sensitive enough reproducible and essential to my success. Thus I find that an appreciation of the pulmonary

megakaryocyte phenomenon and a control of heparin with the Dale and Laidlaw Coagulometer makes heparin prophylaxis understandable and as in my on going study eminently successful.

I will agree with Dr O'Brien that we have much to learn about the actual factors which lead to thrombosis. This I am sure will become evident when we discover just what is derived from our fresh platelets which may suddenly be liberated from the pulmonary megakaryocytes to produce the hypercoagulable state and thrombosis.

J G Sharnoff MD
Pathologist

The Mount Vernon Hospital
Mount Vernon N Y 10550

REFERENCES

- 1 Sharnoff J G, Kass H H and Mistica B A A plan of heparinization of the surgical patient to prevent postoperative thromboembolism *Surg Gynecol Obstet* 115:75 1962
- 2 Sharnoff J G and DeBlasio G Prevention of fatal postoperative thromboembolism by heparin prophylaxis *Lancet* 2:1006 1970
- 3 Kalkar V V, Field E S, Nicolaides A V, Flute P T, Weisler S and Yin H T Low doses of heparin in prevention of deep vein thrombosis *Lancet* 2:669 1971
- 4 Williams H T Prevention of postoperative deep-vein thrombosis with perioperative subcutaneous heparin *Lancet* 2:950 1971
- 5 Gordon Smith I C, Grundy D J, LeQueane L P, Newcombe J F and Bramble F J Controlled trial of two regimens of subcutaneous heparin in prevention of postoperative deep-vein thrombosis *Lancet* 1:1133 1972
- 6 Nicolaides A N, Dupont P A, Desai S, Lewis J D, Douglas J N, Dodsworth H, Fouldes G, Luck H J and Jamieson C W Small doses of subcutaneous sodium heparin in preventing deep vein thrombosis after major surgery *Lancet* 2:890 1972
- 7 Kalkar V V, Corrigan T, Spindler J, Fossard D P, Flute P T, Crellin R Q, Weisler S and Yin E T Efficacy of low doses of heparin in prevention of deep-vein thrombosis after major surgery: a double-blind randomised trial *Lancet* 2:101 1972
- 8 Gallus A S, Hersh J, Tuttle R J, Treblecock R O'Brien S E, Carroll J J, Minden J H and Hudecki S M Small subcutaneous doses of heparin in prevention of venous thrombosis *N Engl J Med* 288:545 1973
- 9 Sharnoff J G and DeBlasio G Some implications in the successful heparin prophylaxis of sudden cardiopulmonary arrest by thrombosis and embolism *Am Heart J* 110:848 1970
- 10 Sharnoff J G Increased pulmonary megakaryocytes—probable role in postoperative thromboembolism *JAMA* 169:688 1973
- 11 Sharnoff J G Prevention of thromboembolism *Bull N Y Acad Med* 49:655 1973
- 12 Sharnoff J G Efficacy of heparin *Lancet* 1:258 1973

Reply

To the Editor

No bibliography is ever complete or satisfies everybody. I apologize for not including a reference to Dr Sharnoff's well known pioneer work with heparin. I omitted this because it is mentioned in the references I quoted and because even though his studies refer to clinical disease I felt the recent work using Iodine 125 fibrinogen in a series of properly controlled studies gives greater precision and makes an even better case for the use of low/small dose heparin—providing that the incidence of clinical disease runs parallel to the incidence of thrombosis diagnosed by this test.

Personally I would not recommend the modified Dale and Ludlaw coagulation time nor do I think there is universal acceptance that the rationale of

this treatment is based on the prevention of a hypercoagulable state due to the liberation of platelets from pulmonary megakaryocytes. But these are matters of opinion.

In the meantime another valuable study supporting the beneficial effects of heparin (and quoting Dr Sharnoff's papers) has recently appeared.¹

J R O'Brien M.A. D.V.
Portsmouth and Isle of Wight Area Pathology Service
Portsmouth England

REFERENCE

- 1 Gallus A S, Hirsh J, Tuttle R J, Trebilcock R, O'Brien S E, Carroll J J, Minden J H and Hudecki S M. Small subcutaneous doses of heparin in prevention of venous thrombosis. *N Engl J Med* 288:545 1973.

Book reviews

CLINICAL PATHOLOGIC CORRELATIONS 2 Jesse E Edwards Guest Editor **CARDIOVASCULAR CLINICS** Albert N. Bresn Editor Philadelphia 1973 F. A. Davis Company 331 pages

The publications of Cardiovascular Clinics continue to be of high quality. They are all clinically oriented and therefore should interest practicing physicians. This issue on the CPC is of considerable value to those who wish to learn the relationship of pathology to the clinical manifestation of heart disease. The diseases presented of broad interest reflect the efforts of authorities in cardiology. Such illnesses as hypertension, ischemic congenital cardiomyopathy and surgical forms of heart disease are discussed. Each chapter is presented as a clinical paper with excellent illustrations and bibliography. This is another excellent contribution to cardiology. Jesse Edwards has produced a useful book for the practicing physician.

ADVANCES IN MICROCIRCULATION Vol 5 H. Harders Editor Basel 1973 B. Karger AG 106 pages

This is another excellent publication on the microcirculation. Volume 5 of *Advances in Microcirculation* is concerned with qualitative and quantitative analysis in blood rheology, intravascular and extravascular morphologic phenomenon in cat skeletal muscle, low flow states and the micro-

circulation of wound healing. These are all extremely important subjects which are discussed clearly and are well illustrated. The authors did not review each field thoroughly; however, they selected aspects of recent advances which they considered most important and interesting. Those who are studying the microcirculation will want to own this publication.

THE UROKINASE PULMONARY EMBOLISM TRIAL American Heart Association Monograph No. 39 Arthur A. Sasahara M.D. Chairman New York 1973 The American Heart Association Inc. 108 pages \$4.00

This paperback publication of a special issue of the American Heart Association summarizes studies of the value of urokinase in the management of pulmonary embolism. The design and results of the study should be primarily of research interest. This complex investigation resulted in supporting the benefit of anticoagulant therapy for patients with pulmonary embolism in the prevention of recurrence. This reviewer is of the strong opinion that an ideal anticoagulant or fibrinolytic is yet to be produced. Those interested in urokinase studies will find the report to be worthy of careful evaluation. Subscribers to *Circulation* already own a copy as a supplement to their subscription to the journal.

Books received

GALAXIES OF LIFE Edited by Stanley Knipper and Daniel Rubin New York 1973 Gordon and Breach 182 pages

RESEARCH COMMUNICATIONS IN CHEMICAL PATHOLOGY AND PHARMACOLOGY Vol 5 Suppl. 1 January 1973 Edited by Richard J. Jones M.D. New York 1973 PJD Publications Ltd. 84 pages

HERZINFARAT KORONARTHROMBOSE UND AKUTER KORONARTOD DES MENSCHEN By Franz Buchner München Berlin Wien 1973 Urban und Schwarzenberg 178 pages

Announcements

Symposium on nutrition

The University of Texas Health Science Center at Houston Division of Continuing Education and the Texas Medical Center Inc. announce the Texas Medical Center's Symposium on the Application of Nutrition in the Health Sciences to be held in Houston Texas on February 8 and 9 1974. Participating in the symposium will be known investigators in the field of nutrition.

For further information please write The Office of the Director The University of Texas Health Science Center at Houston Division of Continuing Education P O Box 20367 Houston Texas 77025.

Conference on cardiac diagnosis and therapy

The Oklahoma Heart Association Colorado Heart Association and the Colorado Society for Cardiovascular Medicine are sponsoring a conference en-

titled 'Dilemmas in Cardiac Diagnosis and Therapy—Circa 1974' to be held at Snowmass Resort Aspen Colorado on March 21 through 23 1974. Featured speakers will be Stephen Epstein M D, Chief of Cardiology Section National Heart Institute Robert J Hall M D Medical Director Texas Heart Institute and W Proctor Harvey M D Professor of Medicine Georgetown University School of Medicine. Also on the program are a distinguished faculty of Oklahoma and Colorado physicians in a series of six sessions emphasizing current problems in the diagnosis and management of common cardiovascular diseases. Registration fee for the conference is \$75.00. A variety of lodge and condominium accommodations are available. Deadline for housing reservations is January 1 1974. Early application is advisable because of limited registration. For further information contact Colorado Heart Association 1375 Delaware Street Denver Colorado 80204 telephone (303) 623 3217.

Author index*

A

- ABELSON DENIS AND MULLER HANS R Reciprocal movement of the right and left heart demonstrated by directional Doppler ultrasound 831
- ABLOQUIST R P Isoproterenol in cardiology 149
- ADAM MASOOD (See Varghese et al) 203
- ALBERT FRANK (See Hatt et al) 805
- ALEMAN JUAN (See de la Torre et al) 467
- AMBAGSI CHRISTOPHER (See Starr et al) 644
- ANDERSON ANTONY K P C (See Stroobandt Fagard and Amery) 181
- AMORI DAVID W (See For 3th et al) 88
- ANBE DANIEL T AND FINE GERALD Cardiac lymphangioma and lipoma Report of a case of simultaneous occurrence in a association with lipomatous infiltration of the myocardium and cardiac arrhythmia 227
- ANDERSON GARY J WOODBURN ROBERT AND FLEMING CHARLES Cerebrovascular accident with unusual electrocardiographic changes 195
- ANDERSON GEORGE A (See de la Torre et al) 467
- ANDRADE ZILTON A AND TEIXEIRA ANTONIO R L Changes in the coronary vasculature in endomyocardial fibrosis and their possible significance 157
- ARCHIE JOSEPH I JR (See Fuxler et al) 188
- ASSISTANT TO THE DIRECTOR FOR MEDICAL COMMUNICATIONS FOOD AND DRUG ADMINISTRATION Digoxin tablet A possible problem with biological availability 715 (Letter to Editor)
- ATTE FALSE (See Horowitz et al) 759
- ATSUMA TAKESHIKO (See Nakayama et al) 96

B

- BARBER HENDRICK B Blood flow in the internal mammary arteries 50 (Annot)
- BAKRESI VINCENT SERRANO ARMANDO COLANDREA MICHAEL A BOGDANOFF MAURICE L AND MLENSTER JOSEPH J Congenital absence of the circumflex coronary artery Clinical and cineangiographic observations 811
- BARON KENNETH (See L Bernstein et al) 817
- BASTA LYNN I (See Zimmerman Basta and Janowitz) 66
- BECKER LEWIS C KLEAS ANDREW F AND HLM PHILIPS J O NEAL Early systolic notch in the aortic arch of man in mitral stenosis 582
- BEVERLY D G BROWN J J FERRIS J B FRANK R LYNN J AND ROBERTSON J L S The use of pronolol in the treatment of hypertension associated with mineralocorticoid excess 404
- BELENKIE ISRAEL CARR MATTHEW SCHLANT R C MITTER D O AND SIMPSON P A Malfunction of a Cutter-Simmons mitral ball valve prosthesis. Diagnosis by phonocardiography and echocardiography 399
- BENAIM RICHARD (See Hatt et al) 285
- BENCHIMOL ALBERTO (See Flahenfeld Desser and Benchimol) 53

- BERGAMASCHI V CARAVAGGI A M MANDELLI V AND SHANKS R G The role of beta adrenoceptors in the coronary and systemic hemodynamic responses to emotional stress in conscious dogs 216
- AND LONGONI ANNA M Cardiovascular events in anxiety. Experimental studies in the conscious dog 185
- BERMAN REuben (See Simonson and Berman) 117
- BHARATI SAROJA AND LEV MAURICE The spectrum of common atrioventricular orifice (canal) 553
- BLAIR DONALD C (See Carr Jr et al) 631
- BLOUNT S GILBERT JR (See Lindquist Spangler and Blount Jr) 286
- BOBB GUSTAVUS A (See Varghese et al) 203
- BODDOVOFF MAURICE L (See Barten et al) 811
- BORDIA ARUN (See Kishori Bordia and Gupta) 282
- BOSSAERT LEO (See Van Durme et al) 284
- BOTTIGER L E AND CARLSON L A Hemoglobin plasma lipids and coronary heart disease 842 (Annot)
- BOWERS DORANCE CHATCOFF VITTE Tooth disease Wolf Parkinson White syndrome and abnormal intracardiac conduction 535
- BREWER A C (See Somerville et al) 822
- BRILLER STANLEY J (See Langer Jr Gewolowitz and Briller) 308
- BRODY DANIEL A (See Copekind et al) 42
- BROOK C G D Cell growth in man 571 (Annot)
- BROWER R W KRALSS V H AND MEESTER G T Level of the base of the mitral valve 745
- BROWN J J (See Beevers et al) 404
- BRUSCA ANTONIO AND POSETTANI ERENIO Activation of the human fetal heart 19
- BRUYNEEL KOENRAAD J J AND OPIE LIONEL H The value of warning arrhythmias in the prediction of entricular fibrillation within one hour of coronary occlusion. Experimental studies in the baboon 373
- BURCH G F Signs of cardiac arrest 138 (Annot)
- 1 chemocardiomyopathy 276 (Annot)
- Pork and hypertension 713 (Annot)
- Study of man himself 475 (Annot)
- Surgical versus medical management of coronary heart disease 846 (Annot)
- AND CALLES T O Ventricular considerations of the post-systolic dip in constrictive pericarditis 569 (Annot)
- AND HARRIS J M Enterohemorrhagic enteritis (EHEC) virus infection of the mouse aorta. An ultrastructural study 669
- AND HIRAHARA Y AND SHIMIZU LANA Viral infection of the aorta of man associated with early atherosclerotic changes 523
- BURCHELL HOWARD B (See Hegge Tuna and Burchell) 603
- BUTTERFIELD W J H (See O'Brien and Butterfield) 711
- BUTTERWORTH J SCOTT AND GLASSMAN EPHRAIM Whether electrocardiography? 709 (Annot)

C

- CACRES C A (See Hochberg et al) 764
 CACIN NORMAN (See Kunzstrdt et al) 173
 CAMERON J S (See Oke and Cameron) 577
 CANJIA ILCIY (See Jacob et al) 438
 CARACTA ANTHONY R (See Josephson et al) 771
 CARAVACCI A M (See Bergman et al) 216
 CARLSON I A (See Bottiger and Carlson) 842
 CARR EDWARD A JR CARROLL MARY DICILLIO WALTER AND BLAIR DONALD C The use of radio-iodinated toluidine blue for myocardial scintigrams 631
 CARR MATTHEW (See Belenkie et al) 399
 CARROLL MARY (See Carr Jr et al) 631
 CASANOVA GÓMEZ M (See Quero-Jiménez et al) 449
 CASTRO GISSONI C (See Quero-Jiménez et al) 449
 CHABOT MICHEL (See Jacob et al) 438
 CHADDA KUT D (See Liebschtein Chadda and Gupta) 13
 CHANG SONIA (See Dillon et al) 698
 CHICHT PAUL (See Huit et al) 285
 CHIEN CHIN CHIN (See Ori cello Chien and Mhdarini) 143
 CHULSKI ANTHONY A ILLIAN PATRICK II AND HILLIS HARRY K The relation ship of coronary collateral inlet flow and retrograde flow in mongrel dogs 485
 COHEN H (See Simon et al) 539
 COHEN KATHI (See Irobst et al) 516
 COHEN MICHAEL V AND ELDBI PER Experimental myocardial infarction in the closed chest dog. Controlled production of large or small areas of necrosis 798
 COHEN SANFORD N DOYLE EUGENE I AND RUTKOWSKI MONIKA M Drug therapy of heart disease in pediatric patients I Congestive heart failure in infancy and concepts of development of pharmacology 133
 — (See Rutkow ki Cohen and Doyle) 270
 — (See Rutkowski Doyle and Cohen) 562
 COLANDREA MICHAEL A (See Barresi et al) 811
 COLLINS ROBERT D (See Butterwhite et al) 107
 COLLARD G DANIEL McLEACHMAN ANN B SMITH HERBERT W AND BRODY DANIEL A The McLeachman (xaxil) vectorcardiogram in normal subjects 42
 CRAIG L ERNST (See Millward McLaughlin and Craig) 848
 CRISTAL N St indard monitor leads—Are reliable as conventional leads? 138 (Annot)

D

- DAUFITT WILLARD M (See Iberthson et al) 817
 DAMATO ANTHONY N (See Vinyhe et al) 203
 — (See Josephson et al) 771
 DAVID DAVID S Dietary treatment of renal failure 1
 DAVIS RICHARD II (See Dillon et al) 698
 DIA TORRE ANGL JACOBS DANIEL ALI MAN JUAN AND ANDERSON GIORI A Embolic coronary artery occlusion in percutaneous transvenous coronary arteriography 467
 DEMOLLEN J C AND KELLYRIS H I Left hemiblock revisited from the histopathological viewpoint 712 (Annot)
 DEMUTH WILLIAM J JR (See Liedtke and De Muth Jr) 687
 DISSER KENNETH B (See Lihensfeld De er and Benchemol) 754
 DHINGRA RAMSII C ROSEN KENNETH M AND RAHIMTOOLA SHAHJAHAN H Wenckebach periods with repetitive block. Evaluation with His bundle recording 443

- DIGILLIO WALTER (See Carr Jr et al) 631
 DILLON JAMES C ILLIENBAUM HARVEY KONECKE III I DAVIS RICHARD II AND CHANG SONIA I electrocardiographic manifestations of valvular vegetations 698
 DINSMORE ROBERT F (See Iberthson et al) 817
 DODER ARTHUR KASCHAUM DONALD G AND GRISWOLD HERBERT E Stress electrocardiography in the evaluation of aortic coronary bypass surgery 292
 DONOSO LEONARDO (See Lyski Stimmel and Donoso) 663
 DOYLE EUGENE I (See Cohen Doyle and Rutkowski) 133
 — (See Rutkow ki Cohen and Doyle) 270
 — (See Rutkowski Doyle and Cohen) 562

F

- EDWARDS I HENRY JR (See Forsyth et al) 88
 ELLIS I N (See Viscoli Iffendy and Maling) 426
 ELDBI PER (See Cohen and Elldbi) 798
 ELLIS KENT (See Steeg Ellis and Gersony) 341
 ELMFELDT DAN (See Vedin et al) 124
 ESINO VITA JORGE (See Horwitz et al) 759
 ESQUIVIA JORGE (See Horwitz et al) 759
 LAIR KENNETH M Complications of trans femoral coronary arteriography and their prevention using heparin 428 (Annot)

F

- FADALI A MONIM A AND SOLOFF LOUIS A An evaluation of human cardiac transplantation 721
 FACARD ROBERT (See Stroobandt Fagard and Amery) 781
 FINKELBAUM HARVEY (See Dillon et al) 698
 FERNANDEZ PAZ (See Kunzstrdt et al) 173
 FERRISS J B (See Beevers et al) 404
 FINE GERALD (See Anbe and Fine) 227
 FISCH CHARLES AND ZILIS DOUGLAS P His bundle electrocardiography 289
 — (See Anderson Woodburn and Fisch) 395
 FISHER JACO DISSER KENNETH B AND BENCHEMOL ALBERTO Non paroxysmal A V junctional tachycardia associated with acute myocardial infarction 754
 FISCHER ERNA Renal excretion of sulfadimidine in normal and uremic subjects 280 (Annot)
 FISLER DAVID L ARCHIBI JOSIEH P JR ULLIOTT DANIEL J AND HOFFMAN JULIAN I Regional coronary flow with increased right ventricular output in anesthetized dog 788
 FLOWER R J A pain like drugs and prostaglandins 814 (Annot)
 FLOWERS NANCY C AND HORAN JIO G Comparative surface potential patterns in obstructive and nonobstructive cardiomyopathy 196
 FORTYTH RAJIN P EDWARDS I HENRY JR AMORY DAVID W MILSON KENNETH I AND THOMSON EAT D Hemodynamic changes during complete heart block in the anesthetized monkey 88
 FOWLER NOBLE O (See Kerner He and Fowler) 625
 FRASER R (See Beevers et al) 404
 FRIEDBERG DAVID I (See Kellon Gillen and Friedberg) 6
 FRIEDMAN SANDOR A PANDYA MAHENDRA AND GRIN LEONST Peripheral arterial occlusion in patients with acute coronary heart disease 415

FRIESINGER GOTTLIEB C (See Safterwhite et al)

107

FILMAN SEYMOUR Therapeutic uses of atrial pacing 833

G

GADGIL R K (See Kapoor et al) 334

GADGIL JOHN J (See Josephson et al) 771

GALLER WILLIAM J (See Helso Callen and Friedberg) 6

GAMBEL OLIVE (See Probst et al) 516

GEORGE M E D (See Hochberg et al) 764

GERSON WELTON M (See Steeg Elh and Gerson) 341

GESLOWITZ DAVID B (See Langner Jr Ceselowitz and Briller) 308

GLASSER IRA H (See Victorica Miller and Gessner) 713

GILES T D (See Burch and Giles) 569

GLASER JORAM Echocardiographic studies of mitral valve 847 (Letter to Editor)

GLASSER STEPHEN P (See Venessee McCarty and Glasser) 57

GLASSMAN EPHRAIM (See Butterworth and Glassman) 109

GLICK G (See Simon et al) 509

GOLDIN BASIL WOLF ELIANA TZIVONI DAN AND STEVEN WILSON Transient S-T elevation detected by 24 hour ECG monitoring during normal daily activity 501

GOLDFREY BRUCE N Reply 280 (Letter to Editor)

GOLDSTEIN SIDNEY (See Zaroff and Goldstein) 681

GOTSMAN M S (See Lewis and Gotzman) 23

GREENBLATT DAVID I AND KOCH WESEY JAN Adverse reactions to propranolol in hospitalized medical patients: A report from the Boston Collaborative Drug Surveillance Program 478

GREENFIELD JOSEPH C JR (See Kendall Rembert and Greenfield Jr) 359

GREIF ERNST (See Friedman Pandya and Greif) 413

GRISWOLD HERBERT E (See Dodek Hassebaum and Griswold) 291

GUNNAR ROLF M (See Karachorlu Gunnar and Arnlower) 238

GUPTA O P (See Kothari Bordia and Gupta) 282

GUPTA PREM K (See Lichstein Chadda and Gupta) 13

GUZON ARTHUR C AND JONES CARL E Central aortic pressure Physiological significance and clinical implication 431

H

HART JACOB I KRANZ PAUL D ALBERT FRANK AND OESTRICHEN ROLF P Protection against epinephrine induced myocardial necrosis with clofibrate 805

HAIAT ROBERT SEBAN CLAUDE BENJAMIN RICHARD AND CHICHE FAYE Right atrial electrocardiogram resulting in hemopericardium 285 (Letter to Editor)

HAMILTON JOHN T (See Khan Hamilton and Manning) 347

HAMNER NINA (See Rubler Schneebism and Hamner) 182

HARR J M (See Burch et al) 523

— (See Burch and Harb) 609

HARRIS WILLARD S (See Pouget and Harris) 286

HARRISON LURA A (See Wittig Harrison and Walke) 69

HARTH KIL J WALKER (See Liberton et al) 817

HASHIBA MITSUAKI (See Matsuo et al) 828

HASHIGUCHI YUO (See Lemura et al) 616

HAYAKAWA H (See Mandel Lozano and Hayakawa) 285

HEATH DONALD (See Somerville et al) 822

HEGGE FREDERICK N TENA NAIR AND BURCHELL HOWARD B Coronary arteriographic findings in patients with axis shifts or S-T segment elevations on exercise stress testing 603

HELLEMS HARPER K (See Cibulki Lehan and Hellem) 485

HENDERSON ROBERT R AND ROWE GEORGE G The progression of coronary atherosclerotic disease as assessed by cine coronary arteriography 165

HENRY M (See Pernot et al) 462

HESS EVELYN V (See Kutsner Hess and Fowler) 675

HIRANOTO Y (See Burch et al) 523

HOCHBERG H M GEORGE M E D SCHMALZ BACH F L AND CACERES C V Automatic ECG and blood pressure measurement in multitest Correlation of blood pressure and ECG abnormalities 764

HOEFFEL J C (See Pernot et al) 462

HOFFMAN JULIEN E (See Fixler et al) 788

HORAN LEO C (See Flower and Horan) 196

HORWITZ SIMON ESQUIVEL A JOSE ARTE FALSA LEM H ELLO AND ESPINOVELA JORCE Clinical diagnosis of per stent left superior vena cava by observation of jugular pulses 759

HULTGREN HERBERT V (See Takaro et al) 587

HUMPHRIES J O NEAL (See Becker Klaus and Humphries) 582

HUNTER JANE (See Probst et al) 516

I

JACKNER I (See Stein et al) 474

JACOB KIRIAN CHABOT MICHEL SALTIEL JACQUES AND CAMPEAU LUCIEN Improvement of left ventricular asynergy following aortocoronary bypass surgery related to preoperative electrocardiogram and vector cardiogram 438

JACOBS DANIEL (See de la Torre et al) 467

JAMES THOMAS V (See Urthaler et al) 189

JANLAKS LEWIS E (See Zimmerman Basta and January) 676

JENZER HANS Reply 282 (Letter to Editor)

JERESATY ROBERT M (See Liss Jeresaty and Vakhouli) 143

JONES CARL E (See Guyton and Jones) 431

JORGENSEN LEIF (See Svendsen and Jorgensen) 144

JOSEPHSON MARK F CARACTA ANTHONY R LAL SEN H GALLAGHER JOHN J AND DAMATO ANTHONY V Electrophysiological evaluation of disopyramide in man 771

J

KAITMAN ALFRED J Indications for aortocoronary artery bypass surgery 420

— Complications of aortocoronary artery bypass surgery 105

KANEHISA TAKUYA (See Lemura et al) 616

KANTER A (See Simon et al) 539

KAPOOR O I MASCARENHAS E RAVANAWARE M M AND GADGIL R K Tuberculosis of the heart Report of 9 cases 334

KARACHORLU ROLF M GUNNAR ROLF M AND ARNLOWER CECIL A Clinical pathologic conference 235

KARNEY JAMES V Experience with the coronary artery bypass graft in a community hospital 51

- SHEEHAN GEORGE A Longevity of athletes 425 (Annot)
- SHELBURNE JAMES C (See Starr et al) 644
- SHEWEY LANA (See Burch et al) 523
- SIEGEL NORMAN J Natural history of childhood lipid nephrosis 139 (Annot)
- SILTZBACH L E (See Stein et al) 474
- SIMON N COHEN H GLICK G KANTLER A LILVIN B AND PIRANI C L Clinical pathologic conference 539
- SIMONSON ERNST AND BERMAN RUBEN New approach in treatment of cardiac decompensation in U S S R 117
- SIVERTSEN EGIL AND JØRGENSEN LIFLIF Reply 144
- SMITH HERBERT W (See Copeland et al) 42
- SOLOFF LOUIS A (See Fadali and Soloff) 721
- SOMERVILLE JANE KHALIL S U BREWER A C AND HEATH DONALD Clinical pathologic conference 822
- SØRLAND SVEIN J (See Rasmussen and Sjørlund) 318
- SOUTHWICK EDWARD G (See Martin Southwick and Muirich) 236
- SPANGLER RICHARD D (See Lindquist Spangler and Blount Jr) 286
- SPODICK DAVID H (See Nandi and Spodick) 495
- STARR ISAAC AMBROSI CHRISTOPHER MANCHESTER JOEL H AND SILLBURN JAMES C Disturbed blood flow in the carotid artery Its physiological and clinical significance 644
- STEEG CARL N ELLIS KENT AND GILSON WELTON M Total anomalous pulmonary venous drainage with ventricular septal defect 341
- STEIN E JACKLER I STIMMEL II STILIN W AND SILTZBACH L E Asymptomatic electrocardiographic alterations in sarcoidosis 474
- STEIN W (See Stein et al) 474
- STERN SHLOMO (See Golding et al) 501
- STIMMEL BARRY (See Stein et al) 474
(See Lipshitz Stimmel and Donato) 663
- STROOBANDT ROLAND IAGARD ROBERT AND ANERY ANTON K P C Are patients with essential hypertension and low renin protected against stroke and heart attack? 781
- SUSMANO ARMANDO (See Barreca et al) 811
- SYMBAS P N (See Belenkie et al) 399

T

- TAKARO TIMOTHY HULTGREN HERBERT N LITT MARY DAVID AND WRIGHT ELIZABETH C An analysis of deaths occurring in association with coronary arteriography 587
- TANAKA HIROMITSU (See Uemura et al) 616
- TELIVIRA ANTONIO R L (See Andrade and Teixeira) 152
- TERASHI SHINICHI (See Uemura et al) 616
- THOMSON PATE D (See Forsyth et al) 888
- TIBBLIN GOSTA (See Vedin et al) 124
- TUNA NAIP (See Hegge Tuna and Burchell) 603
- IZIVONI DAN (See Golding et al) 501

U

- UEMURA NOBUHIKO TANAKA HIROMITSU NIIMURA TATSURU HASHIGUCHI NOBLO YOSHIMURA MASAHITO TERASHI SHINICHI AND KANEHISA TAKUYA Electrophysiological and histological abnormalities of the heart in myotonic dystrophy 616
- ULLYOT DANIEL J (See Fixler et al) 788
- URTHALER FERDINAND KATHOLI CHARLES R MACY JOSIAH JR AND JAMES THOMAS N

Mathematical relationship between automaticity of the sinus node and the AV junction 189

V

- VAN DURM JEAN PIERRE BOSSAERT LEO VERMEIRE PAUL AND PANNIER RENÉ Practical in treating tachyarrhythmias 281 (Letter to Editor)
- VARGHESE P JACON DAMATO ANTHONY N LAU SUN H AKHTAR MASOOD AND BOBB GUS TAVUS A The effect of heart rate acetylcholine and vagal stimulation on antegrade and retrograde His Purkinje conduction in the intact heart 203
- VLADIN J ANDERS WILHELMSSON CLAES ELMFELDT DAC TIBBLIN GOSTA WILHELMSSON LARS AND WERKÖ LARS Sudden death Identification of high risk groups 124
- VERMEIRE PAUL (See Van Durme et al) 284
- VICTORICA BENJAMIN MILLER B LANA AND GESSNER IRA H Electrocardiogram and vectorcardiogram in ventricular inversion (corrected transposition) 733
- VISIOLI O EFFENDI F N AND MALAGUINO G Dopamine test for the diagnosis of coronary insufficiency 426 (Annot)

W

- WALLACE ANDREW G (See Wittig Harrison and Wallace) 69
- WALSTON ABE II AND KENDALL VI ELGENE Comparison of pulmonary wedge and left atrial pressure in man 159
- WATKINS SILVIA M AND LEWIS ADAM Serum enzyme levels after operation 573 (Annot)
- WISSI ALLEN B (See Mochos et al) 61
- WERKÖ LARS (See Vedin et al) 124
- WILHELMSSON LARS (See Vedin et al) 124
- WILHELMSSON CLAES (See Vedin et al) 124
- WITTIG JOHN HARRISON LURA A AND WALLACE ANDREW G Electrophysiological effects of lidocaine on distal Purkinje fibers of canine heart 69
- WOLF ELIANA (See Golding et al) 501
- WOLFF FREDERICK W (See Lapid and Wolff) 211
- WOLLMAN MICHAEL R AND MILLER DENIS R The nitroblue tetrazolium dye test and infection in the renal patient 277 (Annot)
- WONG MAYLNE Depression of cardiac performance by ethanol unmasked during autonomic blockade 508
- WOODBURN ROBERT (See Anderson Woodburn and Fisch) 395
- WORMS A M (See Pernot et al) 162
- WRIGHT ELIZABETH C (See Takaro et al) 587

Y

- YANO KATSUSUKI (See Matsuo et al) 828
- YOSHIMURA MASAHITO (See Uemura et al) 616
- YOSHIOKA MASARO (See Matsuo et al) 828
- YUCCEL LU YUSUF Z (See Kunstadt et al) 173

Z

- ZAROFF LAWRENCE I AND GOLDSTEIN SIDNEY Mitral disc variance (Harken prosthesis) 681
- ZIMMERMAN THOMAS J BASTA LOTFI L AND JANUARY LEWIS E Spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and rheumatic mitral valve disease Description of three patients 676
- ZIPES DOUGLAS P (See Fisch and Zipes) 289

A

- Abdominal films routine postangiography renal abnormalities discovered by (Meneses McCarty and Glasser) 57
- Aberration atrial investigation of as a cause of altered P wave contour (Probst et al) 516
- Abnormal intracardiac conduction Charcot Marie Tooth disease and Wolff Parkinson White syndrome (Bowers) 535
- Absence congenital of the circumflex coronary artery Clinical and cinearteriographic observations (Barresi et al) 811
- Acetylcholine heart rate and vagal stimulation on antegrade and retrograde His-Purkinje conduction in the intact heart effect of (Varghese et al) 203
- Acknowledgment to reviewers 719
- Adrenoceptors beta in coronary and systemic hemodynamic responses to emotional stress in conscious dogs role of (Bergamaschi et al) 216
- Alopecia propranolol induced (Martin Southwick and Maibach) 236
- Angina action of perhexiline maleate in patients with (Morgans and Rees) 329
- Announcements 148 430 576 718 852
- Anodal stimulation as a cause of pacemaker induced ventricular fibrillation (Preston) 366
- Antidepressants tricyclic and cardiac disease (Fox) 841 (Annot)
- Anxiety cardiovascular events in experimental studies in the conscious dog (Bergamaschi and Longoni) 385
- Aorta mouse encephalomyocarditis (EMC) virus infection of the An ultrastructural study (Burch and Harb) 669
- of man viral infection of associated with early atherosclerotic changes (Burch et al) 523
- Aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome nature of pressure flow studies in man (Hendall Rembert and Greenfield) 359
- Aortocoronary artery bypass surgery complications of (Kaltman) 705
- indications for (Kaltman) 420
- bypass surgery improvement of left ventricular asynergy following related to preoperative electrocardiogram and vectorcardiogram (Jacob et al) 438
- stress electrocardiography in evaluation of (Dodek Kassebaum and Griswold) 292
- Apexcardiogram in mitral stenosis early systolic notch in the (Becker Klaus and Humphries) 582
- Are patients with essential hypertension and low renin protected against stroke and heart attack? (Stroobande Fagard and Amery) 781
- Arrest cardiac a sign of (Burch) 138 (Annot)
- Arrhythmias cardiac and lipomatous infiltration of the myocardium report of a case of simultaneous occurrence in association with Cardiac lymphangioma and lipoma (Anbe and Fine) 227
- mechanisms of (Puck) 249
- early following experimental coronary occlusion in conscious dogs and their modification

Arrhythmias—Cont d

- by beta adrenoceptor blocking drugs (Khan Hamilton and Manning) 347
- ventricular lidocaine in (Luss Jereaty and Nakhoul) 143 (Letter to Editor)
- warning value of in the prediction of ventricular fibrillation within one hour of coronary occlusion Experimental studies in the baboon (Bruynel and Opie) 373
- Arterial fistula traumatic coronary (Liberthson et al) 817
- occlusion peripheral in patients with acute coronary heart disease (Friedman Pandya and Greif) 415
- Arteriographic findings coronary in patients with axis shifts or S-T segment elevations on exercise stress testing (Hegge Tuna and Burchell) 603
- Arteriography cine coronary progression of coronary atherosclerotic disease as assessed by (Henderson and Rowe) 165
- coronary an analysis of deaths occurring in association with (Takaro et al) 587
- percutaneous transluminal coronary embolic coronary artery occlusion in (de la Torre et al) 467
- transfemoral coronary complications of and their prevention using heparin (Eyer) 428 (Annot)
- Artery bypass graft coronary in a community hospital experience with (Karnegis) 51
- surgery aortocoronary complications of (Kaltman) 705
- indications for (Kaltman) 420
- carotid disturbed blood flow in the its physiological and clinical significance (Starr et al) 644
- circumflex coronary congenital absence of the Clinical and cinearteriographic observations (Barresi et al) 811
- internal mammary blood flow in the (Barner) 570 (Annot)
- occlusion embolic coronary in percutaneous transluminal coronary arteriography (de la Torre et al) 467
- proximal coronary relation of microcirculatory thrombosis to thrombus in effect of aspirin dipyridamole and thrombolysis (Moschos et al) 61
- Aspirin dipyridamole and thrombolysis effect of relation of microcirculatory thrombosis to thrombus in proximal coronary artery (Moschos et al) 61
- in the prevention of thrombosis (O'Brien and Butterfield) 711 (Annot)
- Aspirin like drugs and prostaglandins (Flower) 844 (Annot)
- Assessment of human cardiac transplantation an (Fadali and Soloff) 721
- Asymptomatic electrocardiographic alterations in sarcoidosis (Stein et al) 474
- Asynergy left ventricular following aortocoronary bypass surgery related to preoperative electrocardiogram and vectorcardiogram improvement of (Jacob et al) 438
- Atherosclerotic changes early viral infection of the aorta of man associated with (Burch et al) 523
- disease coronary as assessed by cine-coronary

- SHEEHAN GEORGE A Longevity of athletes 425 (Annot)
- SHELburnE JAMES C (See Starr et al) 644
- SHELWEY LANA (See Burch et al) 523
- SEIGEL NORMAN J Natural history of childhood lipid nephro 139 (Annot)
- SILTZBACH L E (See Stein et al) 474
- SIMON N COHEN H GLICK G KAVLER A LEVIN B AND PIRANI C I Clinical pathologic conference 539
- SIMONSON ERNST AND BERMAN RUBIN New approach in treatment of cardiac decompensation in U S S R 117
- SIVERTSEN EGIL AND JØRGENSEN LEIF Reply 144
- SMITH HERBERT W (See Copeland et al) 42
- SOLOFF LOUIS A (See Fridly and Soloff) 721
- SOMERVILLE JANE KHALIQ B U BRUWER A C AND HEATH DONALD Clinical pathologic conference 822
- SØRLAND SVEIN J (See Rasmussen and Sørland) 318
- SOUTHWICK EDWARD G (See Martin Southwick and Mubach) 236
- SPINGLER RICHARD D (See Lindquist Spingler and Blount Jr) 286
- SPODICK DAVID H (See Nandi and Spodick) 495
- STARR ISAAC AMBROSI CHRISTOPHE MANCHESTER JOEL H AND SHELburnE JAMES C Disturbed blood flow in the carotid artery Its physiological and clinical significance 644
- STEEG CARL N FLIS KENT AND GLRSON WILTON M Total anomalous pulmonary venous drainage with ventricular septal defect 341
- STEIN E JACKLER I STIMMEL B STEIN W AND SILTZBACH L E Asymptomatic electrocardiographic alterations in sarcoidosis 474
- STEIN W (See Stein et al) 474
- STERN SHLOMO (See Golding et al) 501
- STIMMEL BARRY (See Stein et al) 474
- (See Lip ki Stimmel and Donoso) 663
- STROOBANDT ROLAND FAGARD ROBERT AND AMERY ANTOON K P C Are patients with essential hypertension and low renin protected against stroke and heart attack? 781
- SUSMANO ARMANDO (See Barress et al) 811
- SYMBAS P N (See Belenkie et al) 399

T

- TAKARO TIMOTHY HULTGREN HERBERT N LITTMAN DAVID AND WRIGHT ELIZABETH C An analysis of deaths occurring in association with coronary arteriography 587
- TANAKA HIROMITSU (See Uemura et al) 616
- TEIXEIRA ANTONIO R L (See Andrade and Teixeira) 152
- TERASHI SHINICHI (See Uemura et al) 616
- THOMSON IATI D (See Forsyth et al) 88
- TIBBLIN GOSTA (See Vedin et al) 124
- TUNA NAIF (See Hegge Tuna and Burchell) 603
- TZIVONI DAN (See Golding et al) 501

U

- UEMURA NOBUHIRO TANAKA HIROMITSU NIIMURA TATSURU HASHIGUCHI NORUO YOSHIMURA MASAHIRO TERASHI SHINICHI AND KANCHISA TAKUYA Electrophysiological and histological abnormalities of the heart in myotonic dystrophy 616
- ULLYOT DANIEL J (See Fixel et al) 788
- URTHALER FERDINAND KATHOLI CHARLES R MACY JOSIAH JR AND JAMES THOMAS N

Mathematical relationship between automaticity of the sinus node and the AV junction 189

V

- VAN DURNE JEAN PIERRE BOSSAERT LEO VERMEIRE PALL AND PAVNIER RENÉ Pricetol in treating tachyarrhythmias 284 (Letter to Editor)
- VARGHESE P JACOB DAMATO ANTHONY N LAU SUN H ASHAK MASOOD AND BOBB GAS TAVUS A The effect of heart rate acetylcholine and vagal stimulation on antegrade and retrograde His-Purkinje conduction in the intact heart 203
- VEDIN J ANDERS WILHELMSSON CLAES ELMFELDT DAC TIBBLIN GOSTA WILHELMSEN LARS AND WERKÖ LARS Sudden death Identification of high risk groups 124
- VERMEIRE PALL (See Van Durne et al) 284
- VICTORIA BENJAMIN E MILLER B LYNN AND GESSNER IRA H Electrocardiogram and vectorcardiogram in ventricular inversion (corrected transposition) 733
- VISIOLI O EFFENDY F N AND MALACINO G Dopamine test for the diagnosis of coronary insufficiency 426 (Annot)

W

- WALLACE ANDREW G (See Wittig Harrison and Wallace) 69
- WALSTON IBE H AND KENDALL M ELCENE Comparison of pulmonary wedge and left atrial pressure in man 159
- WATKINS SYLVIA M AND LEWIS ADAM Serum enzyme levels after operation 573 (Annot)
- WELSL ALLEN B (See Mocho et al) 61
- WERKÖ LARS (See Vedin et al) 124
- WILHELMSEN LARS (See Vedin et al) 124
- WILHELMSSON CLAES (See Vedin et al) 124
- WITTIG JOHN HARRISON LURA A AND WALLACE ANDREW G Electrophysiological effects of lidocaine on distal Purkinje fibers of canine heart 69
- WOLF ELIANA (See Golding et al) 501
- WOLFF FREDERICK W (See Lapid and Wolff) 211
- WOLLMAN MICHAEL R AND MILLER DENIS R The nitroblue tetrazolium dye test and infection in the renal patient 277 (Annot)
- WONG MALENE Depression of cardiac performance by ethanol unmasked during autonomic blockade 508
- WOODBURN ROBERT (See Ander on Woodburn and Frisch) 395
- WORMS A M (See Pernot et al) 462
- WRIGHT ELIZABETH C (See Takaro et al) 587

Y

- YANO KATLUSKE (See Matsuo et al) 828
- YOSHIMURA MASAHIRO (See Uemura et al) 616
- YOSHIOKA MASATO (See Matsuo et al) 828
- YUCIOGLU YUSUF Z (See Kunstadt et al) 173

Z

- ZAROFF LAWRENCE I AND GOLDSTEIN SIDNEY Mitral disc variance (Harken prosthesis) 681
- ZIMMERMAN THOMAS J BASTA LOTFY L AND JANUARY LEWIS E Spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and rheumatic mitral valve disease Description of three patients 676
- ZIPES DOUGLAS P (See Fisch and Zipes) 289

Subject index*

A

- Abdominal films routine postangiography renal abnormalities discovered by (Meneses McCarty and Glasser) 57
- Aberation atrial investigation of as a cause of altered P wave contour (Irobo et al) 516
- Abnormal intracardiac conduction Charcot Marie Tooth disease and Wolff Parkinson White syndrome (Bowers) 535
- Absence con ental of the circumflex coronary artery Clinical and cinearteriographic observations (Barresi et al) 811
- Acetylcholine heart rate and vagal stimulation on antegrade and retrograde His-Purkinje conduction in the intact heart effect of (Varhees et al) 203
- Acknowledgment to reviewers 719
- Adrenoceptors beta in coronary and systemic hemodynamic responses to emotional stress in conscious dogs role of (Bergamachi et al) 216
- Allopica propranolol induced (Martin Southwick and Vabach) 236
- Angina action of perhexiline maleate in patients with (Morgans and Rees) 329
- Announcement 148 430 576 718 852
- Anodal stimulation as a cause of pacemaker induced ventricular fibrillation (Preston) 366
- Antidepressants tricyclic and cardiac disease (Mour) 841 (Annot)
- Anxiety cardiovascular events in experimental studies in the conscious dog (Bergamachi and Longoni) 385
- Aorta mouse encephalomyocarditis (EMC) virus infection of the An ultrastructural study (Burch and Harb) 669
- of man viral infection of associated with early atherosclerotic changes (Burch et al) 523
- Aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome nature of pressure flow studies in man (Hendall Rembert and Greenfield) 359
- Aortocoronary artery bypass surgery complications of (Haltman) 405
- indications for (Haltman) 420
- bypass surgery improvement of left ventricular asynergy following related to preoperative electrocardiogram and vectorcardiogram (Jacob et al) 438
- tress electrocardiography in evaluation of (Dodek Kassebaum and Griswold) 292
- Apecardiogram in mitral stenosis early systolic notch in the (Becker Klaus and Humphries) 582
- Are patients with essential hypertension and low renin protected against stroke and heart attack? (Stroobandt Fagard and Amery) 781
- Arrest cardiac signs of (Burch) 138 (Annot)
- Arrhythmias cardiac and lipomatous infiltration of the myocardium report of a case of simultaneous occurrence in association with Cardiac lymphoma and lipoma (Anbe and Fine) 227
- mechanisms of (Pick) 249
- early following experimental coronary occlusion in conscious dogs and their modification

Arrhythmias—Cont d

- by beta adrenoceptor blocking drugs (Khan Hamilton and Manning) 347
- ventricular lidocaine in (Lis Jerecsy and Vakhoul) 143 (Letter to Editor)
- warning value of in the prediction of ventricular fibrillation within one hour of coronary occlusion Experimental studies in the baboon (Brayneel and Opie) 373
- Arterial fistula traumatic coronary (Liberthson et al) 817
- occlusion peripheral in patients with acute coronary heart disease (Friedman Landya and Cress) 415
- Arteriographic finding coronary in patients with axis shifts or S-T segment elevations on exercise-stress testing (Hlegge Tuna and Burchell) 603
- Arteriography cine-coronary progression of coronary atherosclerotic disease as assessed by (Henderson and Rowe) 165
- coronary an analysis of deaths occurring in association with (Takaro et al) 587
- percutaneous transfemoral coronary embolic coronary artery occlusion in (de la Torre et al) 467
- transfemoral coronary complications of and their prevention using heparin (Eyer) 428 (Annot)
- Artery bypass graft coronary in a community hospital experience with (Harnegis) 51
- surgery aortocoronary complications of (Haltman) 705
- indications for (Haltman) 420
- carotid disturbed blood flow in the its physiological and clinical significance (Starr et al) 644
- circumflex coronary congenital absence of the Clinical and cinearteriographic observations (Barresi et al) 811
- internal mammary blood flow in the (Barner) 510 (Annot)
- occlusion embolic coronary in percutaneous transfemoral coronary arteriography (de la Torre et al) 467
- proximal coronary relation of microcirculatory thrombosis to thrombus in effect of aspirin dipyridamole and thrombolysis (Moschos et al) 61
- Aspirin dipyridamole and thrombolysis effect of relation of microcirculatory thrombosis to thrombus in proximal coronary artery (Moschos et al) 61
- in the prevention of thrombosis (O'Brien and Butterfield) 711 (Annot)
- Aspirin like drugs and prostaglandins (Flower) 844 (Annot)
- Assessment of human cardiac transplantation an (Fadali and Soloff) 721
- Asymptomatic electrocardiographic alterations in sarcoidosis (Stein et al) 474
- Asynergy left ventricular following aortocoronary bypass surgery related to preoperative electrocardiogram and vectorcardiogram improvement of (Jacob et al) 438
- Atherosclerotic changes early viral infection of the aorta of man associated with (Burch et al) 523
- disease coronary as assessed by cine coronary

Atherosclerotic—Cont d

- arteriography progression of (Henderson and Rowe) 165
- Athletes longevity of (Sheehan) 425 (Annot)
- Atrial aberration investigation of as a cause of altered P wave contour (Probst et al) 516
- dissociation (Orisicello Chien and Mädarang) 143 (Letter to Editor)
- Reply (Sivertsen and Jørgensen) 144
- ectopic tachycardia (Mandel Luzzo and Haya) 285 (Letter to Editor)
- Reply (Goldreyer) 286
- electrocardiogram right resulting in hemopericardium (Hauat et al) 285 (Letter to Editor)
- fibrillation chronic and rheumatic mitral valve disease spontaneous return of sinus rhythm in older patients with (Zimmerman Basti and January) 676
- level left to-right shunt at after rupture of papillary muscle from acute myocardial infarction (Nagel Ronan and Roberts) 112
- pacing therapeutic uses of (Furman) 835
- pressure left and pulmonary wedge in man comparison of (Walston and Kendall) 159
- Atroventricular orifice (canal) common the spectrum of (Bharati and Lev) 553
- Atropine in acute myocardial infarction (Scherf) 284 (Letter to Editor)
- Automatic ECG and blood pressure measurement in multitest correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- Automaticity of the sinus node and the AV junction mathematical relationship between (Urthaler et al) 189
- Autonomic blockade depression of cardiac performance by ethanol unmasked during (Wong) 508
- AV junction mathematical relationship between automaticity of the sinus node and the (Urthaler et al) 189
- junctional tachycardia non paroxysmal associated with acute myocardial infarction (Fishenfeld Dessar and Benichou) 754
- Avulsed papillary muscle tip infection of simulating bacterial endocarditis (Satterwhite et al) 107
- (Axial) vectorcardiogram in normal subjects the McFee Parungao (Copeland et al) 42
- Axis deviation left complete right bundle branch block with significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- shifts or S T segment elevations on exercise stress testing coronary arteriographic findings in patients with (Hegge Tuna and Burchell) 603

B

- Back syndrome straight Clinical and hemodynamic study of 9 cases (Matsuo et al) 828
- Bacterial endocarditis infection of an avulsed papillary muscle tip simulating (Satterwhite et al) 107
- Ball valve prosthesis Cutter Smeloff mitral mal function of diagnosis by phonocardiography and echocardiography (Belenkie et al) 399
- Beta adrenoceptor(s) blocking drugs early arrhythmias following experimental coronary occlusion in conscious dogs and their

Beta adrenoceptor(s)—Cont d

- modification by (Khan Hamilton and Manning) 347
- in coronary and systemic hemodynamic responses to emotional stress in conscious dogs role of (Bergamaschi et al) 216
- Bifascicular block a clinical and electrophysiologic study (Kunstadt et al) 173
- Biological availability a possible problem with digoxin tablets (Assistant to the Director for Medical Communications Food and Drug Administration) 715 (Letter to Editor)
- Block bifascicular a clinical and electrophysiologic study (Kunstadt et al) 173
- complete heart in patients with LAD RBBB magnitude of risk of developing (Kulbertus) 278 (Annot)
- in the unanesthetized monkey hemodynamic changes during (Forsyth et al) 88
- right bundle branch with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- repetitive Wenckebach periods with evaluation with His bundle recording (Dhingra Rosen and Rahimtoola) 444
- Blockade autonomic depression of cardiac performance by ethanol unmasked during (Wong) 508
- Blocking drugs beta adrenoceptor early arrhythmias following experimental coronary occlusion in conscious dogs and their modification by (Khan Hamilton and Manning) 347
- Blood flow and oxygen consumption in normal and ischemic areas of myocardium effect of noradrenaline on (Marshall and Parritt) 653
- disturbed in the carotid artery its physiological and clinical significance (Stritt et al) 644
- in the internal mammary artery (Barner) 510 (Annot)
- pressure and ECG abnormalities correlation of automatic ECG and blood pressure measurement in multitest (Hochberg et al) 764
- measurement and automatic ECG in multitest correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- Books received 288 717 851
- Book reviews 146 287 430 575 716 851
- Boston Collaborative Drug Surveillance Program a report from adverse reactions to propranolol in hospitalized medical patients (Greenblatt and Koch Weser) 478
- Branch block complete right bundle with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- Bundle branch block complete right with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- Bypass graft coronary artery in a community hospital experience with (Karnegis) 51
- surgery aortocoronary artery complications of (Kaltman) 705
- indications for (Kaltman) 420
- improvement of left ventricular asynergy following related to preoperative electrocardiogram and vectorcardiogram (Jacob et al) 438

- Bypass surgery aortocoronary—Cont d
stress electrocardiography in evaluation of
(Dodek Hasschaum and Griswold) 292
- C
- Canal common atrioventricular orifice the pec-
trum of (Bharati and Lev) 553
- Canine heart electrophysiological effects of lido-
caine on distal Purkinje fibers of (Wittig
Harrison and Wallace) 69
- Cardiac arrest a sin of (Burch) 138 (Annot.)
arrhythmia () and lipomatous infiltration of the
myocardium report of a case of simul-
taneous occurrence in association with
Cardiac lymphangioma and lipoma
(Anbe and Fine) 227
mechanisms of (Pick) 249
- Decompensation in U.S.S.R. new approach in
treatment of (Simonson and Berman)
117
- Disease and tricyclic antidepressants (Moir) 841
(Annot.)
- Injures nonpenetrating a collective review
(Liedtke and DeMuth) 687
- Lymphangioma and lipoma Report of a case of
simultaneous occurrence in association
with lipomatous infiltration of the myo-
cardium and cardiac arrhythmia (Anbe
and Fine) 227
- Performance depress. on of by ethanol unmashed
during autonomic blockade (Wong) 508
- Transplantation human an assessment of (Fadali
and Soloff) 721
- Cardiology isoproterenol in (Ahlquist) 149
- Cardiomyopathy idiopathic immunologic findings
in a prospective serial study (Hirsner
Hess and Fowler) 625
- Ischemic (Burch) 276 (Annot.)
- Obstructive and nonobstructive comparative sur-
face potential patterns in (Flowers and
Hocan) 196
- Of the left ventricle in childhood radiological
patterns of (Pernot et al.) 462
- Cardiovascular events in anxiety experimental
studies in the conscious dog (Berga-
maschi and Longoni) 385
- Cardioversion after valve replacement (Seizer) 282
(Letter to Editor)
- Reply (Jenzer and Lown) 282
- Carotid artery disturbed blood flow in the its
physiological and clinical significance
(Starr et al.) 644
- First derivati = determination of systolic intervals
utilizing the (Nandi and Spodick) 495
- Catheter displacement pacemaker (Klein) 429
(Letter to Editor)
- Reply (Preston) 479
- Cell growth in man (Brook) 571 (Annot.)
- Central venous pressure physiological significance
and clinical implications (Guyton and
Jones) 431
- Cerebrovascular accident with unusual electro-
cardiographic changes (Anderson Wood-
burn and Fisch) 395
- Charcot Marie-Tooth disease Wolff Parkinson
White syndrome and abnormal intra-
cardiac conduction (Bowers) 535
- Childhood lipod nephrosis natural history of
(Siegel) 139 (Annot.)
- Cinearteriographic and clinical observations Con-
genital absence of the circumflex coronary
artery (Barresi et al.) 811
- Cine-coronary arteriography progress on of coro-
nary atherosclerotic disease as assessed
by (Henderson and Rowe) 165
- Circumflex coronary artery congenital absence of
the Clinical and cinearteriographic ob-
servations (Barresi et al.) 811
- Clinical diagnosis of persistent left superior vena
cava by observation of jugular pulses
(Horwitz et al.) 759
- Pathologic conference (Karachorlu Gunnar and
Kraikower) 238
(Simon et al.) 539
(Somerville et al.) 822
- Clofibrate protection against epinephrine induced
myocardial necrosis with (Haft et al.)
805
- Collateral inlet flow coronary and retrograde
flow in mongrel dog relation ship of
(Cibulski Lchan and Hellem) 485
- Common atrioventricular orifice (canal) the pec-
trum of (Bharati and Le) 553
- Complete heart block in patients with LAD RBBB
magnitude of risk of developing (Kul-
bertus) 278 (Annot.)
- Complications of aortocoronary artery bypass sur-
gery (Kaltman) 705
- Of transfemoral coronary arteriography and their
prevention using heparin (Eyer) 428
(Annot.)
- Conduction abnormal intracardiac Charcot Marie-
Tooth disease and Wolff Parkinson
White syndrome (Bowers) 535
- His-Purkinje antegrade and retrograde in the
intact heart effect of heart rate acetyl-
choline and vagal stimulation on
(Varghese et al.) 203
- Congenital absence of the circumflex coronary ar-
tery Clinical and cinearteriographic
observations (Barresi et al.) 811
- Heart disease critical demography of (Kelso
Gallen and Friedberg) 11
- Congestive heart failure in infancy and concepts of
developmental pharmacology I Drug
therapy of heart disease in pediatric
patients (Cohen Doyle and Rutkowski)
133
- in infants and children with digitalis prepara-
tions treatment of II Drug therapy of
heart disease in pediatric patients
(Rutkowski Cohen and Doyle) 270
- Constrictive pericarditis left ventricular function in
systole and diastole in (Lewis and Gots-
man) 23
- post systolic dip of theoretic considerations
of (Burch and Giles) 569 (Annot.)
- Conventional leads standard monitor leads as reli-
able as? (Cristal) 138 (Annot.)
- Coronary and systemic hemodynamic responses to
emotional stress in conscious dogs role
of beta adrenoceptors in (Bergamaschi et
al.) 216
- arterial fistula traumatic (Liberthson et al.) 817
- arteriographic findings in patients with axis shifts
or S-T segment elevations on exercise-
stress testing (Hegge Tuna and Bur-
chell) 603
- arteriography an analysis of deaths occurring in
association with (Takaro et al.) 587
- percutaneous transfemoral embolic coronary
artery occlusion in (de la Torre et al.)
467
- transfemoral complications of and their pre-
vention using heparin (Eyer) 428
(Annot.)
- artery bypass graft in a community hospital
experience with (Karnegis) 51
- circumflex congenital absence of the Clin

Atherosclerotic—Cont d

- arteriography progression of (Henderson and Rowe) 165
- Athletes longevity of (Sheehan) 425 (Annot)
- Atrial aberration investigation of as a cause of altered P wave contour (Probst et al) 516
- dissociation (Orsicello Chien and Mădărang) 143 (Letter to Editor)
- Reply (Sivertsen and Jørgensen) 144
- ectopic tachycardia (Mandel Lozano and Hrya kawa) 285 (Letter to Editor)
- Reply (Goldreyer) 286
- electrocardiogram right resulting in hemopericardium (Haat et al) 285 (Letter to Editor)
- fibrillation chronic and rheumatic mitral valve disease spontaneous return of sinus rhythm in older patients with (Zimmerman Basta and Januury) 676
- level left to-right shunt at after rupture of papillary muscle from acute myocardial infarction (Nagel Roman and Roberts) 112
- pacing therapeutic uses of (Furman) 835
- pressure left and pulmonary wedge in man comparison of (Walston and Kendall) 159
- Atroventricular orifice (canal) common the spectrum of (Bharati and Lev) 553
- Atropine in acute myocardial infarction (Scherf) 284 (Letter to Editor)
- Automatic ECG and blood pressure measurement in multitesting correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- Automaticity of the sinus node and the AV junction mathematical relationship between (Urthaler et al) 189
- Autonomic blockade depression of cardiac performance by ethanol unmasked during (Wong) 508
- AV junction mathematical relationship between automaticity of the sinus node and the (Urthaler et al) 189
- junctional tachycardia non paroxysmal associated with acute myocardial infarction (Fishenfeld Dessur and Benchemol) 754
- Avulsed papillary muscle tip infection of simulating bacterial endocarditis (Satterwhite et al) 107
- (Axial) vectorcardiogram in normal subjects the (McFee Parungao (Copeland et al) 42
- Axis deviation left complete right bundle branch block with significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- shifts or S-T segment elevations on exercise stress testing coronary arteriographic findings in patients with (Hegge Tuna and Burchell) 603

B

- Back syndrome straight clinical and hemodynamic study of 9 cases (Matsuo et al) 828
- Bacterial endocarditis infection of an avulsed papillary muscle tip simulating (Satterwhite et al) 107
- Ball valve prosthesis Cutter Smeloff mitral mal function of diagnosis by phonocardiography and echocardiography (Belenkie et al) 399
- Beta adrenoceptor(s) blocking drugs early arrhythmias following experimental coronary occlusion in conscious dogs and their

Beta adrenoceptor(s)—Cont d

- modification by (Khan Hamilton and Manning) 347
- in coronary and systemic hemodynamic responses to emotional stress in conscious dogs role of (Bergamini et al) 216
- Bifascicular block a clinical and electrophysiologic study (Kunstadt et al) 173
- Biological availability a possible problem with digoxin tablets (Assistant to the Director for Medical Communications Food and Drug Administration) 715 (Letter to Editor)
- Block bifascicular a clinical and electrophysiologic study (Kunstadt et al) 173
- complete heart in patients with LAD RBBB magnitude of risk of developing (Kulbertus) 278 (Annot)
- in the unanesthetized monkey hemodynamic changes during (Forsyth et al) 88
- right bundle branch with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- repetitive Wenckebach periods with evaluation with His bundle recording (Dhingra Rosen and Rahimtoola) 444
- Blockade autonomic depression of cardiac performance by ethanol unmasked during (Wong) 508
- Blocking drugs beta adrenoceptor early arrhythmias following experimental coronary occlusion in conscious dogs and their modification by (Khan Hamilton and Manning) 347
- Blood flow and oxygen consumption in normal and ischemic areas of myocardium effect of noradrenaline on (Marshall and Parrott) 653
- disturbed in the carotid artery its physiological and clinical significance (Starr et al) 644
- in the internal mammary artery (Barner) 570 (Annot)
- pressure and ECG abnormalities correlation of automatic ECG and blood pressure measurement in multitesting (Hochberg et al) 764
- measurement and automatic ECG in multitesting correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- Books received 288 717 851
- Book reviews 146 287 430 575 716 851
- Boston Collaborative Drug Surveillance Program a report from adverse reactions to propranolol in hospitalized medical patients (Greenblatt and Koch Weser) 478
- Branch block complete right bundle with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- Bundle branch block complete right with left axis deviation significance of vectorcardiographic morphology (Lichstein Chadda and Gupta) 13
- Bypass graft coronary artery in a community hospital experience with (Karnegis) 51
- surgery aortic coronary artery complications of (Haltman) 705
- indications for (Haltman) 420
- improvement of left ventricular asynergy following related to preoperative electrocardiogram and vectorcardiogram (Jacob et al) 438

- G-Cont'd
- and blood pressure measurement automatic in multitester correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- and vectorcardiogram in ventricular inversion (corrected transposition) (Victoria Miller and Gessner) 133
- preoperative improvement of left ventricular asynergy following aortocoronary bypass surgery related to (Jacob et al) 438
- effect of heroin and multiple drug abuse on (Lipki Stimmel and Donoso) 663
- monitoring 24-hour transient S-T elevation detected by during normal daily activity (Golding et al) 501
- right atrial resulting in hemopericardium (Haaf et al) 285 (Letter in Editor)
- wide band recording of and coronary heart disease (Langner Gevelowitz and Briller) 308
- Echocardiographic manifestations of valvular vegetations (Dillon et al) 698
- studies of mitral valve (Glaser) 847 (Letter to Editor)
- Reply (Millward McLaughlin and Craige) 848
- Echocardiography and phonocardiography diagnosis by malfunction of a Cutler Sneloff mitral ball valve prosthesis (Belenkie et al) 399
- Ectopic tachycardia atrial (Mandel Losano and Hayakawa) 285 (Letter to Editor)
- Reply (Goldreier) 286
- Electrocardiographic alterations in sarcoidosis asymptomatic (Stein et al) 474
- changes cerebrovascular accident with unusual (Anderson Woodburn and Fisch) 395
- findings in single ventricle and related conditions (Quero-Jiménez et al) 449
- Electrocardiography His bundle (Fisch and Zipes) 289
- stress in evaluation of aortocoronary bypass surgery (Dodek Hassebaum and Griswold) 292
- whither? (Butterworth and Glassman) 709 (Annot)
- Electrophysiologic study a clinical and bifascicular block (Kunstadt et al) 173
- Electrophysiological and histological abnormalities of the heart in myotonic dystrophy (Lemura et al) 616
- effects of lidocaine on distal Purkinje fibers of canine heart (Wittig Harrison and Wallace) 69
- evaluation of disopyramide in man (Josephson et al) 771
- Elevation transient S-T detected by 24 hour ECG monitoring during normal daily activity (Golding et al) 501
- Emboic coronary artery occlusion in percutaneous transluminal coronary arteriography (de la Torre et al) 467
- (EMC) encephalomyocarditis virus infection of the mouse aorta An ultrastructural study (Burch and Harb) 669
- Emotional stress in conscious dogs role of beta adrenoreceptors in coronary and systemic hemodynamic responses to (Bergamaschi et al) 216
- Encephalomyocarditis (EMC) virus infection of the mouse aorta An ultrastructural study (Burch and Harb) 669
- Endocarditis bacterial infection of an avulsed papillary muscle tip simulating (Satterwhite et al) 197
- Endomyocardial fibrosis changes in coronary vasculature in and their possible significance (Andrade and Teixeira) 152
- Enzyme level serum after operation (Watkins and Lewis) 573 (Annot)
- Epiaphrine induced myocardial necrosis with clofibrate protection against (Haft et al) 805
- Essential hypertension and low renin are patients with protected against stroke and heart attack? (Stroobandt Esgard and Amery) 81
- Ethanol depression of cardiac performance by unmasked during autonomic blockade (Wong) 508
- Excretion renal of ulfamididine in normal and uremic subjects (Fraser) 280 (Annot)
- Exercise-stress testing coronary arteriographic findings in patients with axis shifts or S-T segment elevations on (Hegge Tuna and Burchell) 603
- F
- Failure congestive heart in infants and children with digitalis preparations treatment of II Drug therapy of heart disease in pediatric patients (Kutkowski Cohen and Doyle) 210
- Fetal heart human activation of the (Brusca and Rovetani) 19
- Fibrillation atrial chronic and rheumatic mitral valve disease spontaneous return of sinus rhythm in older patients with (Zimmerman Basta and January) 676
- ventricular pacemaker induced anodal stimulation as a cause of (Ireston) 366
- within one hour of coronary occlusion value in warning arrhythmias in the prediction of Experimental studies in the baboon (Bruyneel and Opie) 373
- Fibrosis endomyocardial changes in coronary vasculature in and their possible significance (Andrade and Teixeira) 152
- First derivative carotid determination of systolic intervals utilizing the (Nandi and Spodick) 495
- Fistula traumatic coronary arterial (Liberthson et al) 817
- Flow blood and oxygen consumption in normal and ischemic areas of myocardium effect of noradrenaline on (Marshall and Larratt) 633
- in the internal mammary artery (Barner) 570 (Annot)
- coronary collateral inlet and retrograde flow in mongrel dogs relationship of (Cibulski Lehan and Hellem) 485
- disturbed blood in the carotid artery its physiological and clinical significance (Starr et al) 644
- pattern aortic in both valvular mitral insufficiency and the prolapsed mitral valve syndrome nature of pressure flow studies in man (Hendall Rembert and Greenfield) 359
- regional coronary with increased right ventricular output in anesthetized dogs (Fixler et al) 788
- retrograde in mongrel dogs relationship of coronary collateral inlet flow and (Cibulski Lehan and Hellem) 485
- G
- Gallopp sound left ventricular and acute myocardial

Coronary artery—Cont d

- cal and cinearteriographic observations (Barresi et al) 811
- occlusion embolic in percutaneous transfemoral coronary arteriography (de la Torre et al) 467
- proximal relation of microcirculatory thrombosis to thrombus in effect of aspirin, dipyridamole and thrombolysis (Moschos et al) 61
- atherosclerotic disease as assessed by cine coronary arteriography progression of (Henderson and Rowe) 165
- collateral inlet flow and retrograde flow in mongrel dogs relationship of (Cibulski, Lehan and Hellems) 485
- flow regional with increased right ventricular output in anesthetized dogs (Fixler et al) 788
- heart disease acute peripheral arterial occlusion in patients with (Friedman, Pandya and Greif) 415
- plasma lipids and hemoglobin (Bottiger and Carlson) 842 (Annot.)
- surgical versus medical management of (Burch) 846 (Annot.)
- wide band recording of the electrocardiogram and (Langner, Geselowitz and Briller) 308
- insufficiency dopamine test for diagnosis of (Visiohi, Effendy and Malagnino) 426 (Annot.)
- occlusion early arrhythmias following experimental in conscious dogs and their modification by beta adrenoceptor blocking drugs (Khan, Hamilton and Manning) 347
- value of warning arrhythmias in the prediction of ventricular fibrillation within one hour of Experimental studies in the baboon (Bruyneel and Opie) 373
- vasculature in endomyocardial fibrosis and their possible significance changes in (Andrade and Teixeira) 152
- Cutter Smclloff mitral ball valve prosthesis mal function of diagnosis by phonocardiography and echocardiography (Belenkie et al) 399
- Cyclophosphamide and the treatment of the nephrotic syndrome in adults (Ogg and Cameron) 577

D

- Deaths occurring in association with coronary arteriography an analysis of (Takaro et al) 587
- sudden identification of high risk groups (Vedin et al) 124
- Decompensation cardiac in USSR new approach in treatment of (Simonson and Berman) 117
- Demography of critical congenital heart disease (Kelso, Gallen and Friedberg) 6
- Depression of cardiac performance by ethanol unmasked during autonomic blockade (Wong) 508
- Derivative carotid first determination of systolic intervals utilizing the (Nandi and Spodick) 495
- Deviation lead axis complete right bundle branch block with significance of vectorcardiographic morphology (Lichstein, Chadda and Gupta) 13
- Diastole and systole left ventricular function in constrictive pericarditis in (Lewis and Gotsman), 23

- Dietary treatment of renal failure (David) 1
- Digitalis preparations treatment of congestive heart failure in infants and children with II Drug therapy of heart disease in pediatric patients (Rutkowski, Cohen and Doyle) 270
- Digoxin tablets a possible problem with biological availability (Assistant to the Director for Medical Communications Food and Drug Administration) 715 (Letter to Editor)
- Dip post systolic of constrictive pericarditis theoretic considerations of (Burch and Giles) 569 (Annot.)
- Dipyridamole aspirin and thrombolysis effect of relation of microcirculatory thrombosis to thrombus in proximal coronary artery (Moschos et al) 61
- Directional Doppler ultrasound reciprocal movement of the right and left heart demonstrated by (Abelson and Muller) 651
- Disc variance mitral (Harken prosthesis) (Sharma et al) 681
- Disopyramide in man electrophysiological evaluation of (Josephson et al) 771
- Dissociation atrial (Orsello, Chien and Mada-rang) 143 (Letter to Editor)
- Reply (Sivertsen and Jørgensen) 144
- Distress of dying (the Rees) 141 (Annot.)
- Disturbed blood flow in the carotid artery its physiological and clinical significance (Starr et al) 644
- Dopamine test for diagnosis of coronary insufficiency (Visiohi, Effendy and Malagnino) 426 (Annot.)
- Doppler ultrasound directional reciprocal movement of the right and left heart demonstrated by (Abelson and Muller) 651
- Drug(s) abuse multiple effect of heroin and on the electrocardiogram (Lipski, Stimmei and Donoso) 663
- aspirin like and prostaglandins (Flower) 844 (Annot.)
- failure in reducing pressor effect of isometric handgrip stress test in hypertension (Larind and Wolff) 211
- Surveillance Program Boston Collaborative a report from adverse reactions to propranolol in hospitalized medical patients (Greenblatt and Koch, Weser) 478
- therapy of heart disease in pediatric patients I Congestive heart failure in infancy and concepts of developmental pharmacology (Cohen, Doyle and Rutkowski) 133
- II Treatment of congestive heart failure in infants and children with digitalis preparations (Rutkowski, Cohen and Doyle) 270
- III The therapeutic challenge of supra ventricular tachyarrhythmias in infants and children (Rutkowski, Doyle and Cohen) 562
- Dye test nitroblue tetrazolium and infection in the renal patient (Wollman and Miller) 277 (Annot.)
- Dying the distress of (Rees) 141 (Annot.)
- Dystrophy myotonic electrophysiological and histological abnormalities of the heart in (Uemura et al) 616

E

- ECG abnormalities and blood pressure correlation of automatic ECG and blood pressure measurement in man (Gottlieb et al) 764

- infarction, myocardial acute—Cont'd
- left to-right shunt at atrial level after rupture of papillary muscle from (Nagel Roman and Roberts) 112
 - left ventricular gallop sound and (Riley Russell and Rackley) 598
 - non paroxysmal A V junctional tachycardia associated with (Fishbein, Dessler and Benchemol) 754
 - experimental in the closed chest dog: controlled production of large or small areas of necrosis (Cohen and Eldh) 798
 - infection in the renal patient: nitroblue-tetrazolium dye test and (Wollman and Miller) 277 (Annot.)
 - of involved papillary muscle tip simulating bacterial endocarditis (Satterwhite et al.) 107
 - of the mouse aorta: encephalomyocarditis (F) (VC) virus: An ultrastructural study (Burch and Harb) 669
 - viral of the aorta of man associated with early atherosclerotic changes (Burch et al.) 523
- Inlet flow: coronary collateral and retrograde flow in mongrel dogs: relationship of (Cibulski, Lehan and Hellemis) 485
- Insufficiency: coronary dopamine test for diagnosis of (Visoli, Essendi and Malagnino) 426 (Annot.)
- valvular: mitral: nature of aortic flow pattern in and the prolapsing mitral valve syndrome: pressure-flow studies in man (Kendall, Rembert and Greenfield) 359
- Internal mammary artery: blood flow in the (Barner) 570 (Annot.)
- Intervals: systolic utilizing the carotid first derivative: determination of (Nandi and Spodick) 495
- Intracardiac conduction: abnormal Charcot-Marie-Tooth disease and Wolff-Parkinson-White syndrome (Bowers) 535
- Inversion: ventricular (corrected transposition): electrocardiogram and vectorcardiogram in (Victorica, Miller and Gesner) 733
- Ischemic: areas of myocardium: effect of noradrenaline on blood flow and oxygen consumption in normal and (Marshall and Parratt) 653
- cardiomyopathy (Burch) 276 (Annot.)
- Isometric handgrip stress test in hypertensives on drug failure in reducing pressor effect of (Lamid and Wolff) 211
- Ioprotorenol in cardiology (Ahikist) 149
- J
- Jugular pulses: clinical diagnosis of persistent left superior vena cava by observation of (Horowitz et al.) 759
- Junctional: tachycardia: non paroxysmal A V associated with acute myocardial infarction (Fishbein, Dessler and Benchemol) 754
- L
- LAD RBBB: magnitude of risk of developing complete heart block in patients with (Kulbertus) 278 (Annot.)
- Leads: conventional standard monitor leads as reliable as? (Cristal) 138 (Annot.)
- standard monitor—as reliable as conventional leads? (Cristal) 138 (Annot.)
- Left axis deviation: complete right bundle branch block with significance of vectorcardiographic morphology (Lichstein, Chadda and Gupta) 13
- Left to-right shunt at atrial level after rupture of papillary muscle from acute myocardial infarction (Nagel Roman and Roberts) 112
- Level of base of the mitral valve (Brower, Kraus and Meester) 745
- Lidocaine: electrophysiological effects on distal Lurkin fibers of canine heart (Wittig, Harrison and Wallace) 69
- in ventricular arrhythmia (Liss, Jerecsaty and Vakhoul) 143 (Letter to Editor)
- Lipids: plasma hemoglobin and coronary heart disease (Buttger and Carlson) 847 (Annot.)
- Lipoid nephrosis: childhood: natural history of (Siegel) 139 (Annot.)
- Lipoma and cardiac lymphangioma: Report of a case of simultaneous occurrence in association with lipomatous infiltration of the myocardium and cardiac arrhythmia (Anbe and Fine) 237
- Lipomatous infiltration of the myocardium and cardiac arrhythmia: report of a case of simultaneous occurrence in association with Cardiac lymphangioma and lipoma (Anbe and Fine) 237
- Longevity of athletes (Sheehan) 425 (Annot.)
- Lymphangioma: cardiac and lipoma: Report of a case of simultaneous occurrence in association with lipomatous infiltration of the myocardium and cardiac arrhythmia (Anbe and Fine) 237
- M
- McFee-Parungao (axial) vectorcardiogram in normal subjects (the Copeland et al.) 42
- Maleate: perhexiline action of in patients with angina (Morgans and Rees) 329
- Malfuction of a Cutter Smeloff mitral ball valve prosthesis: diagnosis by phonocardiography and echocardiography (Belenkie et al.) 399
- Mammary artery: internal blood flow in the (Barner) 570 (Annot.)
- Man himself: study of (Burch) 425 (Annot.)
- Mathematical relationship between automaticity of the sinus node and the A-V junction (Urbaher et al.) 189
- Mechanisms of cardiac arrhythmias from hypothesis to physiologic fact (Pick) 249
- Medical versus surgical management of coronary heart disease (Burch) 846 (Annot.)
- Microcirculatory thrombosis: relation of to thrombus in proximal coronary artery: effect of aspirin, dipyridamole and thrombolytics (Moschos et al.) 61
- Mineralocorticoid excess: use of spironolactone in diagnosis and treatment of hypertension associated with (Bevers et al.) 404
- Mitral ball valve prosthesis: Cutter Smeloff: malfunction of: diagnosis by phonocardiography and echocardiography (Belenkie et al.) 399
- disc: variance (Harken prosthesis) (Sharma et al.) 681
- insufficiency: valvular: nature of aortic flow pattern in and the prolapsing mitral valve syndrome: pressure-flow studies in man (Kendall, Rembert and Greenfield) 359
- stenosis: early systolic notch in the apexcardiogram in (Becker, Klaus and Humphries) 582
- valve disease: rheumatic: spontaneous return of sinus rhythm in older patients with

- Gallop sound—Cont d
infarction (Riley, Russell and Rackley) 598
- Graft coronary artery bypass in a community hospital experience with (Karnegis) 51
- II
- Handgrip stress test in hypertension isometric drug failure in reducing pressor effect of (Lamid and Wolff) 211
- Harken prosthesis (mitral disc variance) (Sharma et al.) 681
- Heart attack and stroke are patients with essential hypertension and low renin protected against? (Stroobandt Fagard and Amery) 781
- beat logic claim of voluntary control over an unusual demonstration (Kothari Bordia and Gupta) 282 (Letter to Editor)
- block complete in patients with LAD RBBB magnitude of risk of developing (Kulbertus) 278 (Annot)
- in the unanesthetized monkey hemodynamic changes during (Forsyth et al.) 83
- canine electrophysiological effects of lidocaine on distal Purkinje fibers of (Wittig Harrison and Wallace) 69
- disease coronary acute peripheral arterial occlusion in patients with (Friedman Pandya and Greif) 415
- plasma lipids and hemoglobin (Bottiger and Carlson) 842 (Annot)
- surgical versus medical management of (Burch) 846 (Annot)
- wide band recording of the electrocardiogram and (Langner Geislowitz and Briller) 308
- critical congenital demography of (Kelso Gallen and Friedberg) 6
- in pediatric patients drug therapy of I Congestive heart failure in infancy and concepts of developmental pharmacology (Cohen Doyle and Rutkowski) 133
- II Treatment of congestive heart failure in infants and children with digitalis preparations (Rutkowski Cohen and Doyle) 270
- III The therapeutic challenge of supra-ventricular tachyarrhythmias in infants and children (Rutkowski Doyle and Cohen) 562
- electrophysiological and histological abnormalities of the in myotonic dystrophy (Uemura et al.) 616
- failure congestive in infancy and concepts of developmental pharmacology I Drug therapy of heart disease in pediatric patients (Cohen Doyle and Rutkowski) 133
- in infants and children with digitalis preparations treatment of II Drug therapy of heart disease in pediatric patients (Rutkowski Cohen and Doyle) 270
- human fetal activation of the (Brusca and Rosettani) 79
- intact effect of heart rate acetylcholine and vagal stimulation on antegrade and retrograde His Purkinje conduction in the (Varghese et al.) 203
- rate acetylcholine and vagal stimulation on antegrade and retrograde His Purkinje conduction in the intact heart effect of (Varghese et al.) 203
- right and left reciprocal movement of demonstrated by directional Doppler ultrasound (Abelson and Muller) 651
- Heart—Cont d
tuberculosis of Report of 9 cases (Kapoor et al) 334
- Hemiblock left revisited from the histopathological viewpoint (Demoulin and Kulbertus) 712 (Annot)
- Hemodynamic and clinical study of 9 cases Straight back syndrome (Matsuo et al.) 828
- changes during complete heart block in the unanesthetized monkey (Forsyth et al.) 83
- responses systemic to emotional stress in conscious dogs role of beta adrenoceptors in coronary and (Bergamaschi et al.) 216
- Hemoglobin plasma lipids and coronary heart disease (Bottiger and Carlson) 842 (Annot)
- Hemopericardium right atrial electrocardiogram resulting in (Haat et al.) 285 (Letter to Editor)
- Heparin and venous thrombosis (Sharnoff) 849 (Letter to Editor)
- Reply (O'Brien) 849
- complications of transfemoral coronary arteriography and their prevention using (Eyer) 428 (Annot)
- Heroin and multiple drug abuse on the electrocardiogram effect of (Lipiski Stimmel and Donoso) 663
- High risk groups identification of sudden death (Vedin et al.) 124
- His bundle electrocardiography (Fisch and Zipes) 289
- recording evaluation with Wenckebach periods with repetitive block (Dhingra Rosen and Rahimtoola) 444
- His Purkinje conduction antegrade and retrograde in the intact heart effect of heart rate acetylcholine and vagal stimulation on (Varghese et al.) 203
- Histological and electrophysiological abnormalities of the heart in myotonic dystrophy (Uemura et al.) 616
- Histopathological viewpoint left hemiblock revisited from the (Demoulin and Kulbertus) 712 (Annot)
- Human cardiac transplantation an assessment of (Fadali and Soloff) 721
- fetal heart activation of the (Brusca and Rosettani) 79
- Hypertension and pork (Burch) 713 (Annot)
- drug failure in reducing pressor effect of isometric handgrip stress test in (Lamid and Wolff) 211
- essential and low renin are patients with protected against stroke and heart attack? (Stroobandt Fagard and Amery) 781
- use of spironolactone in diagnosis and treatment of associated with mineralocorticoid excess (Beavers et al.) 404
- I
- Idiopathic cardiomyopathy immunologic findings in a prospective serial study (Kirsner Hess and Fowler) 625
- Immunologic findings in idiopathic cardiomyopathy a prospective serial study (Kirsner Hess and Fowler) 625
- Infancy congestive heart failure in and concepts of developmental pharmacology I Drug therapy of heart disease in pediatric patients (Cohen Doyle and Rutkowski) 133
- Infarction myocardial acute atropine in (Scherf) 284 (Letter to Editor)

- atrial patients—Cont d
 children with digitalis preparations (Rutkowski, Cohen and Doyle) 210
 drug therapy of heart disease in 111 The therapeutic challenge of supraventricular tachyarrhythmias in infants and children (Rutkowski, Doyle and Cohen) 562
 cutaneous transdermal coronary arteriography embolic coronary artery occlusion in (de la Torre et al.) 467
 hexiline maleate action of in patients with angina (Morgans and Rees) 329
 heartitis constrictive left ventricular function in systole and diastole in (Lewis and Gotman) 23
 post systolic dip of theoretic considerations of (Burch and Giles) 569 (Annot)
 peripheral arterial occlusion in patients with acute coronary heart disease (Friedman, Pandya and Grief) 415
 pharmacology developmental congestive heart failure in infancy and concepts of 1 Drug therapy of heart disease in pediatric patients (Cohen, Doyle and Rutkowski) 133
 phonocardiography and echocardiography diagnosis by malfunction of a Cutter Smeloff mitral ball valve prosthesis (Belenkie et al.) 399
 plasma lipids hemoglobin and coronary heart disease (Boutiger and Carlson) 842 (Annot)
 risk and hypertension (Burch) 713 (Annot)
 ultrasonography abdominal film renal abnormalities discovered by routine (Meneses, McCarty and Glasser) 57
 postpartum period and pregnancy systolic time intervals in (Rubler, Schneebaum and Hammer) 182
 post-systolic dip of constrictive pericarditis, theoretic considerations of (Burch and Giles) 569 (Annot)
 potential pattern, comparative surface in obstructive and nonobstructive cardiomyopathy (Flowers and Horan) 196
 prolapsing mitral valve syndrome (Van Durme et al.) 284 (Letter to Editor)
 pregnancy and the postpartum period systolic time intervals in (Rubler, Schneebaum and Hammer) 182
 pressor effect of isometric handgrip stress test in hypertension drug failure in reducing (Lamini and Wolff) 211
 pressure blood and ECG abnormalities correlation of automatic ECG and blood pressure measurement in multitesting (Hochberg et al.) 764
 measurement and automatic ECG in multitesting correlation of blood pressure and ECG abnormalities (Hochberg et al.) 164
 venous physiological significance and clinical implications (Guyton and Jones) 431
 flow studies in man the nature of aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome (Kendall, Rembert and Greenfield) 359
 left atrial and pulmonary wedge in man comparison of (Walston and Kendall) 159
 right ventricular systolic in pulmonary stenosis from combined vectorcardiographic data prediction of (Rasmussen and Spirland) 318
 Prolapsing mitral valve syndrome nature of aortic flow pattern in both valvular mitral insufficiency and pressure-flow studies in man (Kendall, Rembert and Greenfield) 359
 Propranolol induced alopecia (Martin, Southwick and Malsbuchi) 236
 in hospitalized medical patient adverse reactions to a report from the Boston Collaborative Drug Surveillance Program (Greenblatt and Koch-Weser) 478
 Prostaglandins and a pain like drugs (Flower) 844 (Annot)
 Prosthesis Harken (mitral disc valvance) (Sharma et al.) 681
 mitral ball valve malfunction of a Cutter Smeloff diagnosis by phonocardiography and echocardiography (Belenkie et al.) 399
 Protection against epinephrine induced myocardial necrosis with clofibrate (Holt et al.) 805
 Proximal coronary artery relation of microcirculatory thrombosis to thrombus in effect of a (irin dipyridamole and thrombolysis (Mochon et al.) 81
 Pulmonary stenosis prediction of right ventricular systolic pressure in from combined vectorcardiographic data (Rasmussen and Spirland) 318
 venous drainage total anomalous with ventricular septal defect (Steeg, Ellis and Gersony) 341
 wedge and left atrial pressure in man comparison of (Walton and Kendall) 159
 Tube(s) jugular clinical diagnosis of persistent left superior vena cava by observation of (Moravitz et al.) 759
 wave volume a theoretical approach to (Naka, Yama et al.) 96
 Purkinje fibers of canine heart distal electrophysiological effects of lidocaine on (Wittig, Harrison and Wallace) 69
 I wave contour altered investigation of atrial aberration as a cause of (Irobo et al.) 516
 R
 Radio-iodinated colindene blue for myocardial scintigrams use of (Carr et al.) 631
 Radiological patterns of obstructive cardiomyopathy of the left ventricle in childhood (Iernot et al.) 462
 RBBB LAD magnitude of risk of developing complete heart block in patients with (Kulbertus) 278 (Annot)
 Reciprocal movement of the right and left heart demonstrated by directional Doppler ultrasound (Velson and Muller) 651
 Recording of the electrocardiogram wide band and coronary heart disease (Langner, Geselowitz and Briller) 308
 Regional coronary flow with increased right ventricular output in anesthetized dogs (Fowler et al.) 788
 Renal abnormalities discovered by routine post angiography abdominal films (Meneses, McCarty and Glasser) 57
 excretion of sulfadiazine in normal and uremic subjects (Fischer) 280 (Annot)
 failure dietary treatment of (David) 1
 patient nitroblue tetrazolium dye test and infection in (Wollman and Miller) 277 (Annot)
 Renin low are patients with essential hypertension and protected against stroke and heart attack? (Strooband, Fagard and Amery) 781

- Mitral valve—Cont d
 chronic atrial fibrillation and (Zimmerman Basta and January) 676
 echocardiographic studies of (Glaser) 847
 (Letter to Editor)
 Reply (Millward McLaurin and Craige) 848
 level of base of (Brower Krauss and Meester) 745
 syndrome prolapsing nature of aortic flow pattern in both valvular mitral insufficiency and pressure flow studies in man (Kendall Rembert and Greenfield) 359
- Monitor leads standard—is reliable vs conventional leads? (Cristal) 138 (Annot)
- Monkey unanesthetized hemodynamic changes during complete heart block in (Forsyth et al.) 88
- Morphology vectorcardiographic significance of complete right bundle branch block with left axis deviation (Lichstein Chadda and Gupta) 13
- Mouse aorta encephalomyocarditis (EMC) virus infection of the An ultrastructural study (Burch and Hurb) 669
- Muscle tip avulsed papillary infection of simulating bacterial endocarditis (Satterwhite et al.) 107
- Myocardial infarction acute atropine in (Scherf) 284 (Letter to Editor)
 left to-right shunt at atrial level after rupture of papillary muscle from (Nagel Ronan and Roberts) 112
 left ventricular gallop sound and (Riley Russell and Rickley) 598
 non paroxysmal A V junctional tachycardia associated with (Fishenfeld Desser and Benchemol) 754
 experimental in the closed chest dog controlled production of large or small areas of necrosis (Cohen and Eldh) 798
 necrosis with clofibrate protection against epinephrine induced (Haft et al.) 805
 scintigrams use of radio iodinated toluidine blue for (Carr et al.) 631
- Myocardium effect of noradrenaline on blood flow and oxygen consumption in normal and ischemic areas of (Marshall and Parratt) 653
 lipomatous infiltration of and cardiac arrhythmia report of a case of simultaneous occurrence in association with Cardiac lymph angioma and lipoma (Anbe and Fine) 227
- Myotonic dystrophy electrophysiological and histological abnormalities of the heart in (Uemura et al.) 616
- N**
- Natural history of childhood lipid nephrosis (Siegel) 139 (Annot)
- Necrosis controlled production of large or small areas of experimental myocardial infarction in the closed chest dog (Cohen and Eldh) 798
 myocardial with clofibrate protection against epinephrine induced (Haft et al.) 805
- Nephrosis childhood lipid natural history of (Siegel) 139 (Annot)
- Nephrotic syndrome in adults cyclophosphamide and the treatment of the (Ogg and Cameron) 577
- New approach in treatment of cardiac decompensation in USSR (Simonson and Berman) 117
- Nitroblue tetrazolium dye test and infection in the renal patient (Wollman and Miller) 277 (Annot)
- Node sinus and the AV junction mathematical relationship between automaticity of the (Urthaler et al.) 189
- Nonobstructive and obstructive cardiomyopathy comparative surface potential patterns in (Flowers and Horan) 196
- Non paroxysmal A V junctional tachycardia associated with acute myocardial infarction (Fishenfeld Desser and Benchemol) 754
- Nonpenetrating cardiac injuries a collective review (Liedtke and DeMuth) 687
- Noradrenaline on blood flow and oxygen consumption in normal and ischemic areas of myocardium effect of (Marshall and Parratt) 653
- O**
- Obstructive and nonobstructive cardiomyopathy comparative surface potential patterns in (Flowers and Horan) 196
 cardiomyopathy of the left ventricle in childhood radiological patterns of (Pernot et al.) 462
- Occlusion coronary early arrhythmias following experimental in conscious dogs and their modification by beta adrenoceptor blocking drugs (Khan Hamilton and Manning) 347
 value of warning arrhythmias in the prediction of ventricular fibrillation within one hour of Experimental studies in the baboon (Bruyneel and Opie) 373
- embolic coronary artery in percutaneous transfemoral coronary arteriography (de la Torre et al.) 467
- peripheral arterial in patients with acute coronary heart disease (Friedman Pandya and Greif) 415
- Operation serum enzyme levels after (Watkins and Lewis) 573 (Annot)
- Orifice common atrioventricular (canal) the spectrum of (Bharati and Lev) 553
- Oxygen consumption and blood flow in normal and ischemic areas of myocardium effect of noradrenaline on (Marshall and Parratt) 653
- P**
- Pacemaker catheter displacement (Klein) 429
 (Letter to Editor)
 induced ventricular fibrillation anodal stimulation is a cause of (Preston) 366
- Pacing atrial therapeutic uses of (Furman) 835
- Papillary muscle rupture of from acute myocardial infarction left to-right shunt at atrial level after (Nagel Ronan and Roberts) 112
 tip avulsed infection of simulating bacterial endocarditis (Satterwhite et al.) 107
- Patients with essential hypertension and low renin are protected against stroke and heart attack? (Stroobandt Fagard and Amery) 781
- Pediatric patients drug therapy of heart disease in I Congestive heart failure in infancy and concepts of developmental pharmacology (Cohen Doyle and Rutkowski) 133
 drug therapy of heart disease in II Treatment of congestive heart failure in infants and

- Pediatric patients—Cont d**
 children with digitalis preparations (Rutkowski, Cohen and Doyle) 240
 drug therapy of heart disease in 111 The therapeutic challenge of supraventricular tachyarrhythmias in infants and children (Rutkowski, Doyle and Cohen) 562
- Percutaneous transluminal coronary arteriography**
 embolic coronary artery occlusion in (de la Torre et al) 467
- Perhexiline maleate** action of in patients with angina (Morgans and Rees) 329
- Pericarditis** constrictive left ventricular function in systole and diastole in (Lewis and Gotsman) 23
 post systolic dip of theoretic considerations of (Burch and Giles) 569 (Annot)
- Peripheral arterial occlusion** in patients with acute coronary heart disease (Friedman, Pandya and Greif) 415
- Pharmacology** developmental congestive heart failure in infancy and concepts of 1 Drug therapy 2 heart disease in pediatric patients (Cohen, Doyle and Rutkowski) 133
- Phonocardiography and echocardiography** diagnosis by malfunction of a Cutter Smeiloff mitral ball valve prosthesis (Belenkic et al) 399
- Plasma lipids** hemoglobin and coronary heart disease (Bottiger and Carlson) 842 (Annot)
- Pork and hypertension** (Burch) 713 (Annot)
- Postangiography abdominal films** renal abnormalities discovered by routine (Meneses, McCarty and Glasser) 57
- Postpartum period and pregnancy** systolic time intervals in (Rubler, Schneebaum and Hammer) 182
- Post systolic dip of constrictive pericarditis** theoretic considerations of (Burch and Giles) 569 (Annot)
- Potential patterns** comparative surface in obstructive and nonobstructive cardiomyopathy (Flowers and Hoan) 196
- Practolol** in treating tachyarrhythmias (Van Durme et al) 284 (Letter to Editor)
- Pregnancy and the postpartum period** systolic time intervals in (Rubler, Schneebaum and Hammer) 182
- Pressor effect of isometric handgrip stress test in hypertension** drug failure in reducing (Larud and Wolff) 211
- Pressure blood and ECG abnormalities** correlation of automatic ECG and blood pressure measurement in multitest (Hochberg et al) 764
 measurement and automatic ECG in multitest correlation of blood pressure and ECG abnormalities (Hochberg et al) 764
- central venous** physiological significance and clinical implications (Guyton and Jones) 431
- flow studies in man** the nature of aortic flow pattern in both valvular mitral insufficiency and the prolapsing mitral valve syndrome (Kendall, Rembert and Greenfield) 359
- left atrial and pulmonary wedge in man** comparison of (Walston and Kendall) 159
- right ventricular systolic** in pulmonary stenosis from combined vectorcardiographic data prediction of (Rasmussen and Sprlind) 318
- Prolapsing mitral valve syndrome** nature of aortic flow pattern in both valvular mitral insufficiency and pressure-flow studies in man (Kendall, Rembert and Greenfield) 359
- Propranolol** induced alopecia (Martin, Southwick and Maibach) 236
 in hospitalized medical patients adverse reactions to a report from the Boston Collaborative Drug Surveillance Program (Greenblatt and Koch-Weser) 418
- Prostaglandins** and a purin like drugs (Flower) 844 (Annot)
- Prostheses** Harken (mitral disc variance) (Sharma et al) 681
 mitral ball valve malfunction of a Cutter Smeiloff diagnosis by phonocardiography and echocardiography (Belenkic et al) 399
- Protection** against epinephrine induced myocardial necrosis with clofibrate (Halt et al) 805
- Proximal coronary artery** relation of microcirculatory thrombosis to thrombus in effect of a purin dipyridamole and thrombolysis (Mochos et al) 61
- Pulmonary stenosis** prediction of right ventricular systolic pressure in from combined vectorcardiographic data (Rasmussen and Sprlind) 318
- venous drainage** total anomalous with ventricular septal defect (Steeg, Ellis and Gersony) 341
 wedge and left atrial pressure in man comparison of (Walston and Kendall) 159
- Pulse(s)** jugular clinical diagnosis of persistent left superior vena cava by observation of (Hofowitz et al) 759
 wave volume a theoretical approach to (Nakayama et al) 96
- Purkinje fibers** of canine heart distal electrophysiological effects of lidocaine on (Wittig, Harrison and Wallate) 69
- P wave contour** altered investigation of atrial aberration as a cause of (Probst et al) 516
- R**
- Rad o-iodinated toluidine blue** for myocardial scintigrams use of (Carr et al) 631
- Radiological patterns** of obstructive cardiomyopathy of the left ventricle in childhood (Fernet et al) 462
- RBBB LAD** magnitude of risk of developing complete heart block in patients with (Kulbertus) 278 (Annot)
- Reciprocal** movement of the right and left heart demonstrated by directional Doppler ultrasound (Abelson and Muller) 651
- Recording** of the electrocardiogram wide band and coronary heart disease (Langner, Geselowitz and Briller) 308
- Regional coronary flow** with increased right ventricular output in anesthetized dogs (Fuxler et al) 788
- Renal abnormalities** discovered by routine post angiography abdominal films (Meneses, McCarty and Glasser) 57
 excretion of sulfadimidine in normal and uremic subjects (Fischer) 280 (Annot)
 failure dietary treatment of (David) 1
 patient autophluorescent tetrazolium dye test and infection in (Mollman and Miller) 277 (Annot)
- Renin** low are patients with essential hypertension and protected against stroke and heart attack? (Stroobandt, Fagard and Amery) 781

- Repetitive block Wenckebach periods with evaluation with His bundle recording (Dhingra Rosen and Rahimtoola) 444
- Retrograde flow in mongrel dogs relationship of coronary collateral inlet flow and (Cibulski Lehan and Hellems) 485
- Rheumatic mitral valve disease spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and (Zimmerman Basta and January) 676
- Rhythm sinus spontaneous return of in older patients with chronic atrial fibrillation and rheumatic mitral valve disease (Zimmerman Basta and January) 676
- Right bundle branch block complete with left axis deviation significance of vector cardiographic morphology (Lichstein Chadda and Gupta) 13
- Risk groups high identification of sudden death (Vedin et al) 124

S

- Sarcoidosis asymptomatic electrocardiographic alterations in (Stein et al) 474
- Scintigrams myocardial use of radio-iodinated toluidine blue for (Carr et al) 631
- Segment elevations S-T or axis shifts on exercise stress testing coronary arteriographic findings in patients with (Hegge Tuna and Burchell) 603
- Septal defect ventricular total anomalous pulmonary venous drainage with (Stegg Ellis and Gersony) 341
- Serum enzyme levels after operation (Watkins and Lewis) 573 (Annot)
- Shunt left to right at atrial level after rupture of papillary muscle from acute myocardial infarction (Nagel Ronan and Roberts) 112
- Sign of cardiac arrest (Burch) 138 (Annot)
- Single ventricle and related conditions electrocardiographic findings in (Quero-Jiménez et al) 449
- Sinus node and the AV junction mathematical relationship between automaticity of the (Urthaler et al) 189
- rhythm spontaneous return of in older patients with chronic atrial fibrillation and rheumatic mitral valve disease Description of three patients (Zimmerman Basta and January) 676
- Spectrum of common atrioventricular orifice (canal) the (Bharati and Lev) 553
- Spironolactone in diagnosis and treatment of hypertension associated with mineralocorticoid excess use of (Beeyers et al) 404
- Spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and rheumatic mitral valve disease Description of three patients (Zimmerman Basta and January) 676
- S-T elevation transient detected by 24 hour ECG monitoring during normal daily activity (Golding et al) 501
- segment elevations or axis shifts on exercise stress testing coronary arteriographic findings in patients with (Hegge Tuna and Burchell) 603
- Standard monitor leads—as reliable as conventional leads? (Cristal) 138 (Annot)
- Stenosis mitral early systolic notch in the apex cardiogram in (Becker Klaus and Humphries) 582
- pulmonary prediction of right ventricular systolic pressure in from combined vectorcardio-

- Stenosis—Cont d
graphic data (Rasmussen and Sjstrand) 318
- Stimulation nodal as a cause of pacemaker induced ventricular fibrillation (Preston) 366
- Straight back syndrome Clinical and hemodynamic study of 9 cases (Matsuo et al) 828
- Stress electrocardiography in evaluation of aorto-coronary bypass surgery (Dodek Kassebaum and Griswold) 292
- emotional in conscious dogs role of beta adrenoreceptors in coronary and systemic hemodynamic responses to (Bergamaschi et al) 216
- test in hypertension isometric handgrip drug failure in reducing pressor effect of (Lamid and Wolff) 211
- Stroke and heart attack are patients with essential hypertension and low renin protected against? (Stroobandt Fagard and Amery) 781
- Study of man himself (Burch) 425 (Annot)
- Sudden death identification of high risk groups (Vedin et al) 124
- Sulfadimidine renal excretion of in normal and uremic subjects (Fischer) 280 (Annot)
- Superior vena cava persistent left clinical diagnosis of by observation of jugular pulses (Horwitz et al) 759
- Supraventricular tachyarrhythmias in infants and children therapeutic challenge of III Drug therapy of heart disease in pediatric patients (Rutkowski Doyle and Cohen) 562
- Surface potential patterns comparative in obstructive and nonobstructive cardiomyopathy (Flowers and Horan) 196
- Surgical versus medical management of coronary heart disease (Burch) 846 (Annot)
- Systemic hemodynamic and coronary responses to emotional stress in conscious dogs role of beta adrenoreceptors in (Bergamaschi et al) 216
- Systole and diastole left ventricular function in in constrictive pericarditis (Lewis and Gottman) 23
- Systolic intervals utilizing the carotid first derivative determination of (Nandi and Spodick) 495
- notch early in the apexcardiogram in mitral stenosis (Becker Klaus and Humphries) 582
- pressure right ventricular in pulmonary stenosis from combined vectorcardiographic data prediction of (Rasmussen and Sjstrand) 318
- time intervals in man (Pouget and Harris) 286 (Letter to Editor)
- Reply (Lundquist Spangler and Blount) 286
- in pregnancy and the postpartum period (Rubler Schneebaum and Hammer) 182

T

- Tachyarrhythmias practolol in treating (Van Durme et al) 284 (Letter to Editor)
- supraventricular in infants and children therapeutic challenge of III Drug therapy of heart disease in pediatric patients (Rutkowski Doyle and Cohen) 562
- Tachycardia atrial ectopic (Mandel Lozano and Hayakawa) 285 (Letter to Editor)
- Reply (Goldrejer) 286

Tachycardia—Cont'd

- functional non-paroxysmal A V associated with acute myocardial infarction (Fienfeld, Desser and Benichou) 754
- Testing exercise-stress, coronary arteriographic findings in patients with axis shifts or S-T segment elevations on (Hegge, Tuna and Burchell) 603
- Theoretic considerations of the post systolic dip of constrictive pericarditis (Burch and Giles) 569 (Annot)
- Theoretical approach to the volume pulse wave (Nakayama et al) 96
- Therapeutic challenge of supraventricular tachyarrhythmias in infants and children 111
- Drug therapy of heart disease in pediatric patients (Rutkowski, Doyle and Cohen) 563
- uses of atrial pacing (Furman) 835
- Therapy drug of heart disease in pediatric patients
- I Congestive heart failure in infancy and concepts of developmental pharmacology (Cohen, Doyle and Rutkowski) 133
 - II The treatment of congestive heart failure in infants and children with digitalis preparations (Rutkowski, Cohen and Doyle) 270
 - III The therapeutic challenge of supraventricular tachyarrhythmias in infants and children (Rutkowski, Doyle and Cohen) 562
- Thrombolysis a purin and dipyridamole effect of relation of microcirculatory thrombosis to thrombus in proximal coronary artery (Vioschos et al) 61
- Thrombosis aspirin in the prevention of (O'Brien and Butterfield) 711 (Annot)
- microcirculatory relation of to thrombus in proximal coronary artery effect of aspirin dipyridamole and thrombolysis (Vioschos et al) 61
- venous and heparin (Sharnoff) 849 (Letter to Editor)
- Reply (O'Brien) 849
- Thrombus in proximal coronary artery relation of microcirculatory thrombosis to effect of a purin dipyridamole and thrombolysis (Vioschos et al) 61
- Time interval in man systolic (Pouget and Harris) 286 (Letter to Editor)
- Reply (Lindquist, Spangler and Blount) 286
- systolic in pregnancy and the postpartum period (Rubler, Schneebaum and Hammer) 182
- Toluidine blue radio-iodinated for myocardial scintigrams use of (Carr et al) 631
- Transfemoral coronary arteriography complications of and their prevention using heparin (Eyer) 428 (Annot)
- percutaneous embolic coronary artery occlusion in (de la Torre et al) 467
- Transient S-T elevation detected by 24 hour ECC monitoring during normal daily activity (Golding et al) 501
- Transplant on human cardiac an assessment of (Fadali and Soloff) 721
- Transposition corrected—electrocardiogram and vectorcardiogram in ventricular inversion (Victorica, Miller and Gessner) 733
- Traumatic coronary arterial fistula A case report and review of the literature (Liberthson et al) 817
- Treatment of cardiac decompensation in USSR

Treatment—Cont'd

- new approach in (Simonson and Beriman) 117
- Tricyclic antidepressants and cardiac disease (Moir) 811 (Annot)
- Tuberculosis of the heart Report of 9 cases (Hapoor et al) 334
- U
- Ultrasound directional Doppler reciprocal movement of the right and left heart demonstrated by (Abelson and Muller) 651
- Ultrastructural study an Encephalomyocarditis (EMC) virus infection of the mouse aorta (Burch and Harb) 669
- Uremic and normal subjects, renal excretion of sulfadimidine in (Fischer) 280 (Annot)
- USSR new approach in treatment of cardiac decompensation in (Simonson and Beriman) 117
- V
- Valvular stimulation heart rate and acetylcholine on antegrade and retrograde His-Purkinje conduction in the intact heart effect of (Varghese et al) 203
- Valve disease rheumatic mitral spontaneous return of sinus rhythm in older patients with chronic atrial fibrillation and (Zimmerman, Basta and January) 676
- mitral echocardiographic studies of (Glaser) 847 (Letter to Editor)
- Reply (Millward, McLaurin and Craig) 848
- level of base of (Brower, Krauss and Meester) 745
- prosthesis ball Cutter Smeloff mitral malfunction of diagnosis by phonocardiography and echocardiography (Belenkie et al) 399
- replacement cardioversion after (Selzer) 282 (Letter to Editor)
- Reply (Jensen and Lown) 282
- syndrome prolapsing mitral nature of aortic flow pattern in both valvular mitral insufficiency and pressure-flow studies in man (Hendall, Rembert and Greenfield) 359
- Valvular mitral insufficiency nature of aortic flow pattern in and the prolapsing mitral valve syndrome pressure flow studies in man (Hendall, Rembert and Greenfield) 359
- vegetations echocardiographic manifestations of (Dillon et al) 698
- Variance mitral disc (Harken prosthesis) (Sharma et al) 681
- Vasculature coronary in endomyocardial fibrosis and their possible significance changes in (Andrade and Teixeira) 152
- Vectorcardiogram and electrocardiogram in ventricular inversion (corrected transposition) (Victorica, Miller and Gessner) 733
- preoperative improvement of left ventricular asymmetry following aortocoronary bypass surgery related to (Jacob et al) 438
- (axial) in normal subjects the McFee-Parungao (Copeland et al) 42
- Vectorcardiographic data prediction of right ventricular systolic pressure in pulmonary stenosis from combined (Rasmussen and Spilard) 318
- morphology significance of complete right bundle branch block with left axis deviation (Lichstein, Chadda and Gupta) 13

- Vegetations valvular echocardiographic manifestations of (Dillon et al.) 698
- Vena cava superior persistent left clinical diagnosis of by observation of jugular pulses (Horwitz et al.) 759
- Venous drainage total anomalous pulmonary with ventricular septal defect (Steeg Ellis and Gersony) 341
- pressure central physiological significance and clinical implications (Guyton and Jones) 431
- thrombosis and heparin (Sharnoff) 849 (Letter to Editor)
- Reply (O'Brien) 849
- Ventricle left obstructive cardiomyopathy of in childhood radiological patterns of (Pernot et al.) 462
- single and related conditions electrocardiographic findings in (Quero-Jiménez et al.) 449
- Ventricular arrhythmia lidocaine in (Liss Jerecsaty and Nakhoul) 143 (Letter to Editor)
- asynergy left following aortocoronary bypass surgery related to preoperative electrocardiogram and vectorcardiogram improvement of (Jacob et al.) 438
- fibrillation pacemaker induced anodal stimulation as a cause of (Preston) 366
- within one hour of coronary occlusion value of warning arrhythmias in the prediction of Experimental studies in the baboon (Bruyneel and Opie) 373
- function left in systole and diastole in constrictive pericarditis (Lewis and Gotsman) 23
- gallop sound left and acute myocardial infarction (Riley Russell and Rackley) 598
- inversion (corrected transposition) electrocardiogram and vectorcardiogram in (Victoria Miller and Gessner) 733
- output increased right in anesthetized dogs regional coronary flow with (Fixler et al.) 788
- septal defect total anomalous pulmonary venous drainage with (Steeg Ellis and Gersony) 341

Ventricular—Cont'd

- systolic pressure right in pulmonary stenosis from combined vectorcardiographic data prediction of (Rasmussen and Sjöstrand) 318
- Viral infection of the aorta of man associated with early atherosclerotic changes (Burch et al.) 523
- Virus infection of the mouse aorta encephalomyocarditis (EMC) An ultrastructural study (Burch and Harb) 669
- Volume pulse wave a theoretical approach to (Nakayama et al.) 96
- Voluntary control over the heart beat Yogi claim of an unusual demonstration (Kothari Bordia and Gupta) 282 (Letter to Editor)
- W
- Warning arrhythmias value of in the prediction of ventricular fibrillation within one hour of coronary occlusion Experimental studies in the baboon (Bruyneel and Opie) 373
- Wave contour altered P investigation of atrial aberration as a cause of (Probst et al.) 516
- volume pulse a theoretical approach to (Nakayama et al.) 96
- Wenckebach periods with repetitive block evaluation with His bundle recording (Dhingra Rosen and Rahimtoola) 444
- Whither electrocardiography? (Butterworth and Glassman) 709 (Annot.)
- Wide band recording of the electrocardiogram and coronary heart disease (Langner Geselowitz and Briller) 308
- Wolff Parkinson White syndrome Charcot Marie-Tooth disease and abnormal intracardiac conduction (Bowers) 535
- Y
- Yogi claim of voluntary control over the heart beat an unusual demonstration (Kothari Bordia and Gupta) 282 (Letter to Editor)

